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(54) Title: HUMAN IMMUNODEFICIENCY VIRUSES	CALIS	ING AIDS IN A NONHIMAN PRIMATE

(57) Abstract

An isolated human immunodeficiency virus (HIV) type 1, having the identifying characteristics of HIV isolate JC and assigned AIDS Reagent Program Catalog Number 3523, was isolated from an HIV-infected chimpanzee that developed AIDS. This chimpanzee represents the first known animal model of HIV-1 induced AIDS. The substantially full-length (infectious) nucleotide sequences of the HIV-1_{JC} and of HIV-1_{NC} molecular clones are provided herein. This HIV-1_{JC} and HIV-1_{NC} isolates are useful for the preparation of recombinant, attenuated and subunit vaccines, as well as for the preparation of challenge stocks. It is also used as a diagnostic reagent in screening for the presence of HIV-1 in biological samples.

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HUMAN IMMUNODEFICIENCY VIRUSES CAUSING AIDS IN A NONHUMAN PRIMATE

CROSS REFERENCE TO RELATED APPLICATIONS

This application claims priority from United States Provisional Patent Application No. 60/050,548, filed June 23, 1997 and from United States Provisional Patent Application No. 60/057,606, filed September 4, 1997.

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BACKGROUND OF THE INVENTION

This invention concerns unique isolates of human immunodeficiency virus type 1 (HIV-1_{JC}) which are highly infectious *in vivo* and produce acquired immune deficiency syndrome (AIDS) in a nonhuman primate.

The human immunodeficiency viruses types 1 and 2 (HIV-1, HIV-2) are retroviruses which have been implicated as the causative agents of acquired immune deficiency syndrome (AIDS) (Barre-Sinoussi et al. [1983] Science 220:868-871). While skepticism about the exact cause of AIDS has arisen (Duesberg [1991] Proc. Natl. Acad. Sci. 88:1575-1579), a large amount of data has accumulated which supports HIV as the biologic agent of this disease. The most convincing evidence includes: mother to child transmission; transmission

via blood transfusion; and transmission via contaminated blood products (Curran et al. [1984] N. Engl. J. Med. 310:69-75). The development of disease in an animal inoculated with HIV-1 would provide confirmatory evidence for the etiology of AIDS. While AIDS-like disease has been recently demonstrated in HIV-2-infected baboons (Barnett et al. [1994] Science 266:642-646), to date, no species, other than humans, has developed AIDS following infection with HIV-1. Thus, the most appropriate animal model for HIV-1 infection currently does not exist. However, an effective alternative animal model has been developed using simian immunodeficiency virus (SIV) infection of macaques. This model has been extremely useful for investigating the pathogenesis of immunodeficiency virus infection and for the testing of vaccines and therapies (Gardner et al. [1994] in *The Retroviruses*, vol. 3 [Levy, ed.] Plenum Press, New York).

Experimental infection of nonhuman primates with human immunodeficiency virus type 1 (HIV-1) has been described for gibbons (Lusso et al. [1988] J. Immunol. 141:2467-2473), pig-tailed macaques (Barre-Sinoussi et al., supra), and chimpanzees (Alter et al. [1984] Science 226:549-552; Fultz et al. [1986] J. Virol. 58:116-124). However, the development of disease (AIDS) has not been documented in these animals or in any animal species infected with HIV-1. For many years, chimpanzees have been a major focus in the development of animal models for HIV-1 infection and therapy. The ability to consistently infect chimpanzees with several HIV-1 subtypes and to reisolate the virus over extended periods (Fultz et al., supra; Johnson et al. [1993] AIDS Res. Hum. Retroviruses 9:375-378) has made chimpanzees useful for testing vaccine candidates (Berman et al. [1990] Nature 345:622-625; Fultz et al. [1992] Science 256:1687-1690; Girard et al. [1995] J. Virol. 69:6239-6248). However, the lack of disease in HIV-1-infected chimpanzees has raised concern over the relevance of these vaccine studies.

The present invention provides HIV-1 isolates that are infectious to nonhuman primates such as chimpanzees and induce AIDS in inoculated nonhuman primates including, but not limited to, chimpanzees. Further, this invention provides for the first time a proper animal model for HIV-1 infection and for the development of AIDS.

SUMMARY OF THE INVENTION

A first aspect of the present invention are the isolated human immunodeficiency virus type 1 (HIV-1) isolate having the identifying characteristics of HIV-1 isolate JC (HIV-1_{JC}) and assigned AIDS Reagent Program Catalog Number 3523 and the isolate having the identifying characteristics of HIV-1 isolate NC (HIV-1_{NC}). These viruses are useful for the preparation of recombinant, attenuated and subunit vaccines, as well as for the preparation of challenge stocks. Each is also useful in screening for the presence of HIV in biological samples.

A second aspect of the invention is a biological sample, e.g., a biological fluid or tissue, containing an HIV-1 having the identifying characteristics of HIV-1_{IC} or HIV-1_{NC}. In a particular embodiment of the invention, primate blood specimens containing the HIV-1_{IC} or the HIV-1_{NC} of the present invention were used to cause HIV infection and to induce AIDS in a nonhuman primate, as specifically exemplified by the chimpanzee.

A third aspect of the invention are a biologically pure culture of host cells containing an HIV-1 having the identifying characteristics of HIV-1_{IC} and a biologically pure culture of host cells containing an HIV-1 having the identifying characteristics of HIV-1_{NC}.

A further aspect of the invention are isolated DNA molecules which produce infectious HIV-1 having the identifying characteristics of HIV-1_{JC} or which encode an antigenic fragment thereof or having the identifying characteristics of HIV-1_{NC} or encode an antigenic fragment thereof. The substantially full length sequence for infectious molecular clone of HIV-1_{JC} is given in SEQ ID NO:11, and the corresponding sequence for the infectious molecular clone of HIV-1_{NC} is given in SEQ ID NO:12.

An additional aspect of the invention provides isolated DNA encoding an HIV-1 envelope protein having the amino acid sequence of SEQ ID NO:2. In a particular embodiment, the DNA sequence encoding an HIV-1 envelope protein contains the nucleotide sequence of SEQ ID NO:1.

A further aspect of the invention are a composition comprising an antigenic preparation derived from the HIV-1_{IC} of the invention and a composition comprising an antigenic preparation derived from the HIV-1_{NC} of the present invention.

An additional aspect of the invention are pharmaceutical compositions comprising an immunogenic amount of an antigenic preparation derived from the HIV-1_{IC} of the invention in a pharmaceutically acceptable carrier or HIV-1_{NC} in a pharmaceutically acceptable carrier.

A further aspect of the present invention is a kit for detecting the presence of HIV-1 antibodies, comprising an antigenic preparation derived from the foregoing HIV-1_{JC} or from HIV-1_{NC} as described herein.

Another aspect of the invention is a method of inducing in a subject antibodies to the HIV-1_{JC} or to HIV-1_{NC} of the invention, comprising the step of administering to the subject an immunogenic amount of an antigenic preparation derived from the HIV-1_{JC} or from the HIV-1_{NC}. The induced antibodies can be harvested and find use, for example, as hybridization probes in methods for HIV scanning.

A further aspect of the invention is a method of immunizing a subject against infection by HIV, comprising the step of administering to the subject an immunogenic amount of an antigenic preparation derived from the HIV-1_{JC} or derived from HIV-1_{NC} of the present invention.

It is another aspect of the invention to provide a method for inducing acquired immune deficiency syndrome (AIDS) in a nonhuman primate, comprising the step of administering to said primate an effective amount of an antigenic preparation derived from the HIV-1_{IC} and/or HIV-1_{NC} of the invention such that said primate develops AIDS. In a specific embodiment of the invention, HIV-1_{IC} induced AIDS in chimpanzees, e.g., in a chimpanzee referred to as C499. In a different embodiment of the invention, HIV-1_{IC} contained in a biological sample from a chimpanzee having AIDS (C499) induces AIDS in a second chimpanzee and in a different primate species.

It is another aspect of the invention to provide a method for inducing acquired immune deficiency syndrome (AIDS) in a nonhuman primate, comprising the step of administering to said primate an effective amount of an antigenic preparation derived from the HIV-1_{NC} of the invention such that said primate develops AIDS. In a specific embodiment of the invention, HIV-1_{NC} induced AIDS in chimpanzees, e.g., in a chimpanzee referred to as C534. In a different embodiment of the invention, HIV-1_{NC} contained in a biological sample from a chimpanzee having AIDS (C455) induces AIDS in a second chimpanzee and in a different primate species.

A further aspect of the invention provides a biological fluid or tissue sample, obtained from a chimpanzee having HIV-1_{JC}-induced AIDS, containing an antigenic HIV fragment for inducing AIDS in a nonhuman primate.

A further aspect of the invention provides a biological fluid or tissue sample, obtained from a chimpanzee having HIV-1_{NC}-induced AIDS, containing an antigenic HIV fragment for inducing AIDS in a nonhuman primate.

It is yet another aspect of the invention to provide a method for inducing AIDS in a nonhuman primate, comprising the step of administering to said primate an effective amount of a biological fluid or tissue sample, obtained from a primate having HIV-1_{JC}-induced or HIV-1_{NC}-induced AIDS, containing an antigenic HIV fragment for inducing AIDS in a nonhuman primate.

The present invention further provides various vaccine formulations containing active immunogenic agents derived from the foregoing HIV-1_{JC}, DNA encoding the HIV-1_{JC}, and DNA encoding antigenic fragments of the HIV-1_{JC} from the foregoing HIV-1_{NC}, DNA encoding the HIV-1_{NC}, and DNA encoding antigenic fragments of the HIV-1_{NC}. An antigenic fragment contains one or more epitopes which bind antibodies directed to the HIV-1_{JC} and/or HIV-1_{NC} of the invention.

BRIEF DESCRIPTION OF THE FIGURES

Fig. 1 shows the CD4+ -cell decline and plasma virus loads in chimpanzee C455 following transfusion with blood from C499. Immediately before and at various times after transfusion, blood was collected from C455 for *in vitro* analyses of CD4+ -cell levels and plasma virus loads. Absolute peripheral CD4+ cells in C455 showed a dramatic decrease by 2 weeks post-transfusion. This rapid decline continued and, by 14 weeks after transfusion, the number of CD4+ cells decreased to 10/µl. These low cell numbers have been maintained to date (42 weeks post-transfusion). Plasma HIV-1 RNA loads in C455 showed high levels of virus present by two weeks after transfusion. Results obtained at the 4-week point suggest very high levels of virus replication in C455. However, HIV-1 levels appeared to be somewhat controlled by 5 to 9 weeks post-transfusion. The cutoff level of 10⁴ equivalents/ml (---) is the lower limit of the assay.

Fig. 2 illustrates CD4+ -cell decline in chimpanzee C534 following transfusion with blood from C455. Immediately before and at various times after transfusion, blood was collected from C534 for *in vitro* analyses of CD4+ -cell levels. Absolute peripheral CD4+ cells in C534 showed a dramatic decrease by 20 days post-transfusion.

Figure 3 illustrates the strategy used in PCR amplification of subgenomic fragments from HIV-1 and the location and orientation of primers on the viral genome are shown. All the primers were designed from the HIV-1_{LAI} nucleotide sequence and their coordinates are described below. PCR amplified fragments were cloned in TA vectors and the corresponding name designations for the recombinant plasmids are shown in bracket. The 5'-LTR-containing Apa I fragment amplified from HIV-1_{IC} PBMC genomic DNA was subcloned in pJC to generate plasmid pHIV-1_{ICI6} while the Apal-Ncol fragment containing the 5'-LTR region amplified from HIV-1_{NC} PBMC genomic DNA was subcloned in pHIV-1_{ICI6} to generate pHIV-1_{NCIC} chimeric plasmid. Plasmid pHIV-1_{NC7} was constructed by subcloning the env-containing Ncol-Xhol fragment amplified from HIV-1_{NC} genomic DNA to pHIV-1_{NCIC} chimeric plasmid. All recombinant plasmids (pHIV-1_{ICI6}, pHIV-1_{NCIC}, and pHIV-1_{NCI}) lacked 55 bp at the 5' end of the genome (U3 region) and all of U5 region in the 3' LTR region.

Figures 4A-4F show replication of cloned and uncloned HIV-1 isolates in con-A stimulated and unstimulated chimpanzee PBMC (cPBMC). Stimulated cPBMC infected with (Fig. 4A) HIV-a LAV-1b and SF2 parental strains and the highly cytopathic DH12 isolate, (Fig. 4B) JC (uncloned) and JC16 (cloned) isolates of HIV-1, and (Fig. 4C) NC (uncloned) and NC7 (cloned) isolates of HIV. Unstimulated cPBMC infected with: (Fig. 4D) LAV-1b, SF2, and DH12; (Fig. 4E) JC, JC16; and (Fig. 4F) NC, NC7. Chimpanzee PBMC (1.1 x 10⁷) in T-25 cm² were infected with 20 ng of either HIV-1_{NC} (uncloned), HIV-1_{SF2}, HIV-1_{LAV-1b}, or HIV-1_{DH12} virus and incubated for a total of 17 days at 37°C. Supernatant aliquots were made on 3, 7, 10, 14, and 17 days post infection. Reverse transcriptase (RT) assays were performed as outlined hereinbelow.

Figure 5 shows replication of HIV-1 isolates in chimpanzee monocyte-derived macrophages (MDM). Purified PBMC (6 x 10⁶/well) were used to obtain MDM. Ten ng of each of virus HIV-1_{IC16} (molecular clone), HIV-1_{IC} (uncloned), HIV-1_{NC7} (molecular clone), HIV-1_{NC} (uncloned), HIV-1_{SF2}, HIV-1_{LAV-1b}, and HIV-1_{DH12}) was used for infection and on days 7 and 14 post infection supernatants were harvested. The mount of virus in the supernatants was determined using the p24 HIV-1 antigen capture ELISA (Coulter). Supernatants from control uninfected cultures are represented.

DETAILED DESCRIPTION OF THE INVENTION

The following definitions are given in order to provide clarity as to the intent or scope of their usage in the specification and claims.

The term HIV-1 isolate JC or HIV-1_{JC} or HIV-1 having the identifying characteristics of HIV-1_{JC} as used herein refers to the particular HIV-1 isolated from a chimpanzee (C499) that developed AIDS 10 years after infection with HIV-1 (Novembre et al. [1997] J. Virol. 71:4086-4091). The DNA sequence of the cloned HIV-1_{JC} is given in SEQ ID NO:11.

The term HIV-1 isolate NC or HIV-1_{NC} or HIV_{NC} or HIV-1 having the identifying characteristics of HIV-1_{NC} as used herein refers to the particular HIV-1 isolated from a chimpanzee (C455) that developed AIDS after infection with HIV-1_{JC} (Novembre et al. [1997] J. Virol. 71:4086-4091). The DNA sequence of the cloned HIV-1_{NC} is given in SEQ ID NO:12.

The term antigenic preparation of HIV-1_{JC} or antigenic fragment of HIV-1_{JC} as used herein refers to the whole viral particle of the HIV-1_{JC} or to a fragment thereof, wherein such fragment encodes at least one epitope or antigenic determinant. The term antigenic preparation of HIV-1_{NC} or antigenic fragment of HIV-1_{NC} as used herein refers to the whole viral particle of the HIV-1_{JC} or to a fragment thereof, wherein such fragment encodes at least one epitope or antigenic determinant.

The term animal or subject as used herein refers to a mammal and, more frequently, to a primate.

The term effective amount as used herein refers to the quantity of active ingredient necessary to effect in an animal a change in a specific biochemical or immunological parameter. For example, in a particular embodiment of this disclosure, an effective amount refers to the amount of HIV-1 administered to a subject such that viral infection and AIDS developed.

The term antigenic amount as used herein refers to the quantity of active antigen necessary to effect an interaction with corresponding antibodies.

The term immunogenic amount as used herein refers to the quantity of active antigen necessary to stimulate the immune system in response to a specific antigen.

The term labeled is used herein to refer to the conjugating or covalent bonding of any suitable detectable group, including enzymes (e.g., horseradish peroxidase, β -glucuronidase, alkaline phosphatase, and β -D-galactosidase), fluorescent labels (e.g., fluorescein, luciferase),

and radiolabels (e.g., ¹⁴C, ¹³¹I, ³H, ³²P, and ³⁵S) to the compound being labeled. Techniques for labeling various compounds, including proteins, peptides and antibodies are well known. See, e.g., Morrison, *Methods in Enzymology* 32B, 103 (1974); Syvanen et al., *J. Biol. Chem.* 284, 3762 [1973]; Bolton and Hunter, *Biochem. J.* 133, 529 [1973].

The viruses of the present invention resulted from the development of AIDS in a first chimpanzee infected with HIV-1 for over 10 years and by the rapid development of immunosuppression in a second chimpanzee transfused with blood from the first chimpanzee. To investigate the ability of HIV-1 to induce AIDS in non-human primates, a cohort of 12 chimpanzees was inoculated with several strains of HIV-1 at the Yerkes Center in the mid-1980s. A member of this cohort, C499, was described as part of a previously reported superinfection study (Fultz et al. [1987] J. Virol. 61:4026-4029) and was inoculated on three separate occasions with three different HIV-1 isolates: HIV-1_{SF2} in 1985, HIV-1_{LAI} in 1986, and HIV-1_{NDK} in 1987. The first inoculation resulted in infection, as determined by positive virus isolation and persistent HIV-1-specific antibody response. Clinically, the animal remained healthy except for the development of thrombocytopenia and lymphopenia in 1988, which resolved without treatment (Fultz et al. [1991] J. Infect. Dis. 163:441-447). (All HIV-infected chimpanzees are maintained in Biosafety level 3 isolation facilities.)

In 1993, after the resumption of yearly monitoring (monitoring of all HIV-1-infected chimpanzees at the Yerkes Center was suspended from March 1990 to May 1993), a decrease in the levels of platelets and CD4⁺ cells in C499 was observed (24,000 and 390/µl, respectively) (Table 1). Thrombocytopenia and CD4⁺-cell lymphopenia were persistent in this animal from this point onward. In addition, C499 displayed other significant clinical signs of disease. Beginning in March 1995, C499 developed chronic, intermittent diarrhea for which no enteric pathogens were identified and which was not resolved with antibiotic treatments. In September 1995, this animal developed acute fulminant diarrhea which was associated with large numbers of *Blastocystis hominis* and *Balantidium coli*. At this time, CD4⁺ cells decreased to extremely low levels (minimum of 10/µl:2% of total T cells) (Table 1), indicative of severe immunosuppression. Similarly, declines were observed in the levels of total lymphocytes and CD8⁺ cells (Table 1). Treatment with fluid replacement and

antimicrobial and antiprotozoal therapy (doxycycline, ceftriaxone, enrofloxacin, gentamicin, and albendazole) resulted in the resolution of acute diarrhea within five days.

Because this was the first chimpanzee to develop AIDS, treatment with antiretroviral therapies was not administered in order to more fully characterize virological, immunological, and pathological parameters in this animal. As the acute diarrhea was resolved with treatment, CD4*-cell levels rose to a maximum of 180/µl (4% of total T cells) but declined again. Subsequently, chronic, intermittent diarrhea resumed and continued unresolved. During this period, C499 exhibited no lymphadenopathy or wasting. However, beginning in the latter part of 1995 and extending into 1996, C499 developed progressive nonregenerative anemia (hematocrit levels of 37.5% in December 1995 and 27.2% in January 1996, with respective hemoglobin values of 12.1 and 8.4 g/dl; reticulocyte counts were 0.0% since November 1995). Due to progressive hematologic abnormalities, chronic diarrhea, and continued immunosuppression, the animal was euthanized in February 1996.

Concomitant with the decrease in CD4⁺ cells was an increase in HIV-1 loads in plasma (Table 1). The increase in the level of the virus was detected in plasma samples dating from May 1993, when the CD4⁺-cell decline was first noted, but not before the suspension of monitoring in 1990. These levels are significantly higher than those for five other chimpanzees at the Yerkes Center which received cell-free or cell-associated HIV-1_{LAI} or HIV-1_{SF2} inoculations (all have undetectable plasma viral RNA levels and are not immunosuppressed). These results suggest that pathogenic effects began to occur sometime between 1990, when CD4⁺-cell counts were at normal levels and viral loads were undetectable, and 1993, when alterations in the number of CD4⁺ cells and significant virus loads were present.

The ability to isolate virus from C499 varied since C499's first exposure to HIV-1. In general, early after inoculation, HIV was easily isolated from the peripheral blood mononuclear cells (PBMC) of this animal; however, after several months, virus could no longer be isolated. From August 1988 until the suspension of monitoring in 1990, the virus was consistently isolated from C499 on a monthly basis. Subsequently, after the resumption

of monitoring in mid-1993, HIV-1 continued to be easily isolated from this animal. Quantitative titration of PBMC viral load following the development of acute diarrhea in C499 in September 1995 until the time of euthanasia revealed that 10⁴ to 10⁵ PBMC was consistently required for virus isolation. The immune response of C499 to HIV-1 infection was very strong up to the time of euthanasia. HIV-1 antibody endpoint titers (HIV-1 whole-virus enzyme-linked immunosorbent assay, Genetic Systems, Redmond, WA) ranged from 51,200 to 204,800 since 1993. Because of the deteriorating condition of this animal and because of the severe decline in CD4⁺ cells, it was hypothesized that the HIV-1 present in this animal had evolved to become more cytopathic for chimpanzee CD4⁺ cells. Cocultivation of PBMC derived from C499 (obtained at the time of acute diarrhea) with uninfected chimpanzee PBMC (cPBMC) resulted in the isolation of a virus (HIV-1_{3C}) which induced syncytium formation in chimpanzee cells (Fig. 1A-D). This characteristic has been previously described for only three HIV-1 isolates (Ghosh et al. [1993] Virology 194:858-864; Schuitemaker et al. [1993] J. Infect. Dis. 168:1140-1147; Shibata et al. [1995] J. Virol. 69:4453-4462), none of which were used for inoculation of C499.

The virus strain derived from C499 (HIV-1_{IC}) and other HIV-1 isolates (HIV-1_{LAI} and HIV-1_{SF2}) were tested for the ability to induce syncytium formation in cPBMC. Virus stocks were prepared in cPBMC (HIV-1_{IC}) or in human PBMC (HIV-1_{LAI} and HIV-1_{SF2}). Cells were incubated with virus overnight and were then washed. Cultures were examined daily for evidence of cytopathic effects. cPBMC four days after infection with an HIV-1 isolate (HIV-1_{IC}) from C499 showed beginnings of syncytium formation and separated syncytia. cPBMC four days after infection with HIV-1_{LAI} lacked of syncytium formation, and normal cell clusters were present. cPBMC four days after infection with HIV-1_{SF2} also showed no syncytium formation. The only virus to induce significant cytopathic effects in cPBMC was HIV-1_{IC}. All cultures were examined for 14 days following infection. All cultures, regardless of the virus used, became positive for virus replication by seven days postinfection. Thus, it is concluded that genetic changes which confer the ability to induce syncytium formation occurred in the virus present in C499.

To confirm that the virus present in C499 was different from the viruses used for inoculation, DNA prepared from HIV-1_{IC}-infected cPBMC was used as a template in typical PCR assays with primers (forward, no. 384:

5'CCCTTCGAAGAGGATATAATCAGTTTATGGGATCAAAGC3' [SEQ ID NO:9]; reverse, no. 383: 5'CCCTTCGAACTCTTCTTCTGCTAGACTGCCATT3' [SEQ ID NO:10]) designed to amplify a 507-bp fragment of the env gene containing the V1 and V2 regions. Genetic analysis of 16 HIV-1_{IC} V1-V2 clones obtained by ligation of the amplification products with the vector pGEM7ZF (Promega, Madison, WI) showed amino acid homologies of 80 to 84% with HIV-1_{LAI}, 73 to 80% with HIV-1_{SF2}, and 63- to 68% with HIV-1_{NDK}. Thus, there appears to be considerable divergence between the virus present in C499 at the time of acute disease and the viruses used to inoculate this animal. This divergence is further illustrated in Fig. Tables 3A-3C, which show amino acid alignments of five HIV-1_{IC} clones, HIV-1_{NDK}, HIV-1_{LAI}, and HIV-1_{SF2}. Comparative analyses between the 16 HIV-1_{JC} clones showed that amino acid homologies ranged from 81 to 96%, with no clones being identical. These results suggest that the virus population in C499 consisted of a large quasispecies. Furthermore, the data, when combined with the in vitro analyses described above, indicate that the virus adapted after years of replication and mutation, becoming more pathogenic for the chimpanzee. While no evidence of recombination is evident from analyses performed in this small area, the possibility of recombination cannot be ruled out for other portions of the HIV-1_{IC} genome.

Tables 2A-2C illustrate amino acid alignment of V1-V2 clones obtained from HIV-1_{JC}-infected cPBMC and prototypes HIV-1_{LAI} and HIV-1_{SF2}, and HIV-1_{NDK}. env clones of HIV-1_{JC} encompassing the V1-V2 region were sequenced by the dideoxy chain termination method (Sequenase; Amersham Life Science, Arlington Heights, IL). With the Intelligenetics Suite of programs (Intelligenetics, Beaverton, OR), sequences of HIV-1_{JC} env fragments were used to derive corresponding amino acid sequences. Deduced amino acid sequences from five of these clones (HIV_{JC}10, HIV_{JC}17, HIV_{JC}45, HIV_{JC}48, and HIV_{JC}55) were then aligned with the corresponding region in HIV-1_{LAI}, HIV-1_{SF2}, and HIV-1_{NDK} isolates. Amino acid sequences of HIV-1_{LAI}, HIV-1_{SF2} and HIV-1_{NDK} were obtained from the Human Retroviruses and AIDS Database.

To investigate additional pathogenic effects of HIV-1 infection on C499, tissue samples obtained by biopsy during the acute diarrheal stage (September 1995) and at necropsy were subjected to histopathological analyses. First, examination of a peripheral lymph node (obtained from C499 in September 1995) revealed marked lymphoid depletion within the cortical area, with a few follicles remaining, as compared with a lymph node from an age-matched, uninfected chimpanzee, which lacked follicular development and had a very cellular cortex. To detect virus expression in tissues, in situ hybridization experiments with digoxigenin-labeled riboprobes encompassing the entire HIV-1 genome (derived from the HIV-1 BH10 molecular clone (Hahn et al. [1984] Nature 312:166-169) were used to probe formalin-fixed lymph node sections. Detection of bound probes was performed with a sheep anti-digoxigenin-alkaline phosphatase-labeled Fab monoclonal antibody and with Nitro Blue Tetrazolium-5-bromo-4-chloro-3-indolylphosphate toluidinium as the substrate chromogen, according to previously described methods (Hirsch et al. [1995] J. Virol. 69:955-967). In situ hybridization studies of C499's lymph node demonstrated the presence of HIV-1 RNA in follicular trapping patterns, with few positively staining cells. These findings are similar to changes observed in HIV-1-infected persons developing AIDS (Pantaleo et al. [1995] N. Engl. J. Med. 332:209-216). Control samples, which included sense riboprobes and lymph nodes from uninfected chimpanzees, did not show any staining. Additional lymph nodes obtained from C499 at necropsy showed a similar depleted pattern. However, a few lymph nodes were not as depleted but did contain multinucleated giant cells which stained positive for the HIV-1 p24 antigen. These giant cells are often found in the lymph nodes of simian immunodeficiency virus (SIV)-infected macaques and are occasionally found in the lymph nodes in HIV-1-infected persons.

Finally, histopathologic analysis of intestinal tissue from C499 revealed pathologic changes in the ileum, with significant blunting of the villi and intense infiltration of mononuclear cells and plasma cells. Examination of the intestinal mucosa at higher magnification revealed extensive infection with *Cryptosporidium*, which was present throughout the small intestine, lining the apical surfaces of intestinal epithelial cells. This organism is an AIDS-defining opportunistic pathogen (Centers for Disease Control and Prevention. 1992, 1993 revised classification system for HIV infection and expanded

surveillance case definition for AIDS among adolescents and adults. Morbid. Mortal. Weekly Rep. 41:1-19) and probably accounted for the chronic intermittent diarrhea and intestinal pathology in C499.

Most tissues obtained from C499 following euthanasia appeared grossly normal, with no lymphadenopathy or splenomegaly. However, the spleen showed moderate congestion and lacked follicle development. Bone marrow specimens obtained at euthanasia showed some functional impairment in CFU granulocyte macrophage and CFU formation, although levels of CD4⁺ cells were not altered (Villinger et al. [1997] J. Med. Primatol 26:181-189. Virus was isolated from all lymphoid organs including inguinal, axillary, and mesenteric lymph nodes as well as from the spleen and thymus. Virus was also isolated from the kidney and liver but not from the brain or cerebrospinal fluid.

At the time of acute diarrhea in C499, a blood transfusion was performed to determine the effects of passage of virus from this animal to an uninfected chimpanzee. Forty milliliters of blood obtained by venipuncture from C499 was immediately transfused intravenously into C455. This chimpanzee, which was bred in captivity, was seropositive for Epstein-Barr virus and cytomegalovirus and was seronegative and PCR negative for HIV prior to the transfusion. Results of titration analysis of PBMC and plasma from C499 show that in the 40 ml of blood, 1 x 10⁴ 50% tissue culture infective doses (TCID₅₀) of virus was in PBMC and 2 x 10⁴ TCID₅₀ of virus was in plasma. Thus, C455 received a total of 3 x 10⁴ TCID₅₀ of virus. Analysis of peripheral CD4⁺ cell levels in C455 revealed a precipitous decline beginning by 2 weeks after transfusion, when absolute numbers of CD4+ cells decreased from 1,240 to 320 cells/µl (Fig. 1). This decline continued, reaching a minimum value of 10 cells/µl (1% of total T cells) by 14 weeks post-transfusion. Since this time, the total percentage of T cells that the CD4⁺ population encompasses has remained constant at 1%, with only a slight rise in the absolute number of CD4⁺ cells (20 cells/µl at the latest time point). Levels of peripheral CD8⁺ cells showed an initial decline from 1,590 to 880 cells/µl in the first two weeks after transfusion. However, these levels quickly rebounded to 2,010 CD8⁺ cells/µl by 8 weeks after transfusion. Since this time, the number of CD8+ cells in circulation has been maintained between 550 and 4,320/µl, with the level in most recent sample being 840 cells/µl.

Because CD4+-cell levels declined so rapidly, the level of virus present in C455 was investigated by quantitation of plasma HIV-1 RNA with the Chiron B-DNA assay (Fig. 1). Two weeks after transfusion, plasma viral loads were 2 x 107 RNA equivalents/ml and reached maximum levels by 4 weeks post-transfusion (6.2 x 10⁷ RNA equivalents/ml). Following a decline (which corresponded with the development of anti-HIV-1 specific antibody), plasma HIV-1 levels have been maintained at ~1.1 x 105 RNA equivalents/ml. Virus has been easily isolated from C455 at all times post-transfusion. Quantitative coculture of PBMC from C455 has shown that early after transfusion, virus could be isolated from as few as 10² PBMC, while at more recent time points, up to 10⁵ to 10⁶ cells was required for virus isolation. In vitro analysis of virus isolated from this animal at several time points has shown that the ability to induce syncytium formation in cPBMC has been retained, further implicating this virus in the pathogenesis of CD4+-cell decline. Antibody responses to HIV-1 have been moderate in C455. Enzyme-linked immunosorbent assay titers have been maintained between 1,600 and 6,400 since four weeks post-transfusion. Clinically, C455 has appeared normal, except for episodic incidences of a rash on the chest and in the scrotum area. In addition, the animal has experienced no weight loss, lymphadenopathy, or anemia.

This invention presents an HIV-1 infected chimpanzee that developed AIDS as defined by the Centers for Disease Control and Prevention classification system (CDCP 1992, supra). Progression of clinical disease (anemia, thrombocytopenia, and chronic diarrhea) in this animal was associated with several key findings, including the following: (i) the presence of a virus which is cytopathic for cPBMC in vitro and in vivo and is genetically distinct from those used for inoculation, (ii) an increase in viral load; (iii) CD4⁺-cell depletion, (iv) lymph node depletion, and (v) the presence of Cryptosporidium organisms in the intestine. It appears that the critical change(s) associated with clinical progression may have developed during the period in which C499 was not monitored (mid-1990 through mid-1993). The precipitous CD4⁺-cell decline concomitant with high viral loads displayed in C455, transfused with blood from C499, suggest that the HIV present in C499 has evolved to become more pathogenic for chimpanzees. The increased pathogenicity of a lentivirus after passage into a new host has been previously observed with the adaptation of SIV from sooty

mangabeys to pig-tailed macaques, resulting in the development of the acutely lethal strain SIVsmmPBi14 (Fultz et al. [1989] AIDS Res. Hum. Retroviruses 5:397-409).

Forty milliliters of blood obtained by venipuncture from C455 was immediately transfused intravenously into C534. This chimpanzee, which was bred in captivity, was seropositive for Epstein-Barr virus and cytomegalovirus and was seronegative and PCR negative for HIV prior to the transfusion. Results of titration analysis of PBMC and plasma from C455 show that in the 40 ml of blood, 1 x 10⁴ 50% tissue culture infective doses (TCID₅₀) of virus was in PBMC and 2 x 10⁴ TCID₅₀ of virus was in plasma. Thus, C534 received a total of 3 x 10⁴ TCID₅₀ of virus. Analysis of peripheral CD4⁺ cell levels in C534 revealed a precipitous decline beginning by one week after transfusion, when absolute numbers of CD4⁺ cells decreased from approximately 1,450 to 700 cells/μl (Fig. 2). The total percentage of T cells that the CD4⁺ population encompasses has remained constant at approximately 1%, with only a slight rise in the absolute number of CD4⁺ cells (20 cells/μl at the latest time point). Levels of peripheral CD8⁺ cells showed an initial decline in the first two weeks after transfusion. However, these levels quickly rebounded after transfusion.

Table 3 shows the antibody response in chimpanzee C534 following transfusion with blood from C455. Immediately before and at various times after transfusion, blood was collected from C534 for *in vitro* analyses of antibody response. Plasma HIV-1 RNA loads in C534 showed high levels of virus present by two weeks after transfusion. Results obtained at the 3-week point indicate very high levels of virus replication in C455. The cutoff level of 10^4 equivalents/ml is the lower limit of the assay.

Because CD4*-cell levels declined so rapidly, the level of virus present in C534 was investigated by quantitation of plasma HIV-1 antibody response (Table 4). One week after transfusion, the plasma antibody titer was still 0. At two weeks after transfusion, the antibody titer rose to 400 and remained at 1600 at three and four weeks post-transfusion. Virus has been easily isolated from C455 at all times after transfusion. Quantitative coculture of PBMC from C534 has shown that early after transfusion, virus could be isolated from as few as 10² PBMC, while at more recent time points, up to 10⁵ to 10⁶ cells was required for virus

isolation. In vitro analysis of virus isolated from this animal at several time points has shown that the ability to induce syncytium formation in cPBMC has been retained, further implicating this virus in the pathogenesis of CD4*-cell decline. Antibody responses to HIV-1 have been moderate in C534. Enzyme-linked immunosorbent assay titers have been maintained between 1,600 and 6,400 since four weeks post-transfusion. Clinically, C534 has appeared normal, except for episodic incidences of a rash on the chest and in the scrotum area. In addition, the animal has experienced no weight loss, lymphadenopathy, or anemia (normal hematocrit levels).

This invention presents HIV-1 isolates NC and JC which infected chimpanzees such that the HIV-1 infected chimpanzees develop AIDS as defined by the Centers for Disease Control and Prevention classification system (CDCP 1992, supra). Progression of clinical disease (anemia, thrombocytopenia, and chronic diarrhea) in this animal is associated with several key findings, including the following: (i) the presence of a virus which is cytopathic for cPBMC in vitro and in vivo and is genetically distinct from that used for inoculation, (ii) an increase in viral load; (iii) CD4⁺-cell depletion, (iv) lymph node depletion, and (v) the presence of Cryptosporidium organisms in the intestine. It is expected that the critical change(s) associated with clinical progression develop during the period spanning approximately 3 to 5 years after HIV-1 infection. The precipitous CD4⁺-cell decline concomitant with high viral loads displayed in C534, transfused with blood from C455, indicates that the HIV present in C455 remains pathogenic for chimpanzees.

In the past, relevance of the HIV-1-infected chimpanzee as a model for vaccine evaluation was questioned due to lack of disease development. The lack of an animal model which supports pathogenic HIV-1 infection has been a continuing problem for vaccine development. Although the time of progression to disease (>10 years), the currently limited numbers of animals available for use, and the overall high costs associated with working with chimpanzees are deterrents to their widespread use in AIDS research, the potential usefulness of this model cannot be disregarded. The development of AIDS in C499, the fact that additional HIV-infected chimpanzees have depressed CD4*-cell counts (<500/µl) and thrombocytopenia, and the rapid progression of the CD4*-cell decline in C455 support the

role that this animal model provides in AIDS-related studies. The adaptation of HIV-1 from long-term chimpanzee infection to a pathogenic form provides a critical link for the adaptation of HIV-1 to growth in more readily available nonhuman primate species. The instant invention further contemplates the growth of HIV-1_{IC} isolated from C499 and/or HIV-1_{NC} from C455 in chimpanzees as well as in pig-tailed or rhesus macaques. In addition, the present invention provides for continued biological and genetic characterization of HIV-1_{IC} and HIV-1_{NC} with further key insights into the pathogenesis of HIV-1 infection in humans and chimpanzees, for example, for the development of drugs and vaccines for the treatment and prevention of AIDS.

As described above, chimpanzee C499 was initially infected with HIV-1_{SF2} in 1985 [Fultz et al. (1986) *J. Virol.* 58, 116-124] and later inoculated with HIV-1_{LAV-1b} and HIV-1_{NDK} in 1986 and 1987 respectively. Superinfection with HIV-1_{LAV-1b} (but not HIV-1_{NDK}) was demonstrated by restriction enzyme analysis of PBMC genomic DNA. At the time of disease development in C499, a virus isolate, termed HIV-1_{IC}, was obtained by co-culture of C499 PBMC with normal cPBMC. At that time, sequence analysis of the V1-V2 region of env suggested that HIV-1_{IC} was most closely related to HIV-1_{LAV}. Also at the time of disease development, blood from C499 was transfused into an uninfected chimpanzee (C455) which resulted in a dramatic decline of CD4+ cells by 2 weeks post transfusion. The depressed CD4+ cell count is still maintained to date in this animal. One month post transfusion 50 μl of plasma from C455 was used for in vitro infection of normal chimpanzee PBMC and the resultant virus was designated HIV-1_{NC}.

To perform a more thorough analysis of the genetic makeup of the HIV-1_{JC} and HIV-1_{NC} viruses, we constructed substantially full-length, infectious molecular clones as described in the Examples and in Figure 3. Both clones lacked 55 nucleotides at the 5' end (5' LTR, U3 region) and all of the U5 region in the 3' LTR. Several clones (representing both HIV-1_{JC} and HIV-1_{NC}) which appeared to be the correct size were tested for biological activity by transfection of CEMx174 cells. Supernatants from transfected cells were used in RT assays to monitor virus production. Two clones, one from each group (HIV-1_{JC} [JC16] and HIV-1_{NC} [NC7]), were positive by RT and also showed massive syncytia formation (2 to 3 days post

transfection), similar to that observed with uncloned virus. To prepare stock viruses for use in in vitro assays, 293 cells were transfected with molecularly cloned DNAs followed by amplification with cPBMC as outlined hereinbelow.

The complete nucleotide sequences of JC16 and NC7 were determined as described in Example 7, and the sequences are given in SEQ ID NO:11 and SEQ ID NO:12, respectively. The genomes were determined to be 9193 nt (JC16) and 9196 nt (NC7) in length and contained open reading (ORFs) for all HIV-1-specific structural, regulatory and accessory genes. Alignment of JC16 and NC7 DNA sequences revealed that the two genomes were very similar, but contained a number of nucleotide changes spread throughout the genome. compared with the parental inoculating viruses, and with gaps introduced to optimize alignment and treated as mismatches, the LTR sequences of JC16 and NC7 had percent nucleotide identities of 92.1% (LAV), 93.9% SF2), and 89.1% (NDK)—lower than that The most divergent region between NC7 and JC16 was the V5 region of the env gene. JC16 contained a 6 bp deletion in the gag gene relative to NC7 while NC7 had 3 bp deletion in the

Comparative analyses were performed between JC16, NC7, and the parental inoculating viruses, SF2, LAV, and NDK (The sequence of LAV-1b has not yet been determined). When observed upon direct comparison of JC16 and NC7 (Table 3). Most of the host/virus transcription binding factor sequences (sites for AP-1, NF-AT, NF-kB and Sp-1) and the TAR CORE and the Lys-tRNA sites were conserved (or had single point mutations) between the parental (SF2 and LAV strains) and progeny viruses. However, there were 3 point mutations unique to JC16 and NC7 at the NRF/NRE binding site.

env gene region relative to JC16. In the LTR region there was a 98.7% nucleotide identity between JC16 and NC7 with all the changes being localized to the U5 region (Table 4).

The deduced amino acid sequences for all proteins of HIV-1_{JC} and HIV-1_{NC} were generated using the Intelligenetics suite of programs. Using the Lasergene program (DNASTAR Inc., Madison, WI), multiple alignments of all proteins were constructed to examine similarities between HIV-1_{NC}, HIV-1_{JC} and the parental viruses (Table 4). Based upon percent homology calculations, Gag, Pol, Vif, Tat, Rev, Env, and Vpu of JC16 and NC7

were most closely related to LAV, with vpr and Nef being most closely related to SF2. In no case was it apparent that the NDK isolate was the origin of a protein sequence. While most changes involved amino acid point mutations, several proteins of JC16 and NC7 contained amino acid insertions or deletions relative to the parental inoculating strains. A closer analysis of amino acid alignments revealed that the percent homologies could be misleading with regards to the origin of the protein. For example, in Tat, JC16 and NC7 were more homologous to LAV than to SF2 (based upon percent homologies). However, the JC16 and NC7 Tat proteins contained 15 amino acid deletions with respect to LAV--similar to that present in the SF2 isolate. Similar findings were observed in Gag and Pol.

Table 5 shows amino acid alignments of Gag, Nef, and Env proteins from JC16, NC7 and the inoculating viruses, LAV, SF2, and NDK. The deduced amino acid sequences for the Gag (a), Nef (b), and Env proteins of HIV-1 were determined using the Intelligenetics Suite of programs (Intelligenetics, Beaverton, OR) and the CLUSTAL method (DNASTAR Inc., Madison, WI) was used for the alignment of proteins. The virus isolates are shown by the designations at the beginning of every line as LAI (HIV-1_{LAV}), JC16 (HIV-1_{JC16}), NC7 (HIV-1_{NC7}), and SF2 (HIV-1_{SF2}). Dashes (-) denote amino acid deletion while dots (.) denote amino acid identity. The numbers after the amino acid sequence on the right show the position of the right-most amino acid in the line. The functional domains on Gag (a) and Nef (b) are indicated. MBD (a) refers to membrane-binding domain while MTD (b) refers to membranetargeting domain. The hypervariable regions in the Env glycoprotein (c) are shown as V1 to V5 and the CD4 binding domain is shown above the sequence. Gp120 (SU) is the outer surface membrane Env glycoprotein. The NDK Env, LAI Env, and SFS Env protein sequences are given in SEQ ID Nos: 3, 4 and 5, respectively. The JC16 and NC7 Env protein sequences are given in SEQ ID Nos: 2 and 23, respectively. There were 7 (JC16 and LAI sequences) and 5 (NC7 sequence) amino acid deletions in the matrix protein (p17) relative to SF2 isolate sequence. At the C-terminal end of Gag polyprotein, the progeny viruses and SF2 virus had 12 amino acid deletions in p6 protein relative to LAV virus sequence (Table 5). However, the capsid (p24) and nucleocapsid (p7) proteins were generally well conserved including the cysteine residues within the zinc-finger domains. Point mutations unique to JC16 and NC7 were present in p17, p24, p7 and p6 peptides. The Lck binding domain

(proline-rich region) within Nef was well conserved with only one point mutation in LAI (Table 5). However, there were 4 amino acid insertions in the SF2 sequence relative to the other viruses at the N-terminal portion of Nef. Sequence analysis of eight other non-infectious clones (4 from JC and 4 from NC) confirmed observations made for the Gag and Nef deletions suggesting that these characteristics are a general property of the viruses obtained from C499 and C455.

The vast majority of point mutations, deletions, and insertions in these clones, relative to the parental viruses, were found in the env glycoprotein region (Tables 4 and 5). Alignment of the Env proteins revealed that all the cysteine residues resident in the protein were conserved between the viruses. There were a total of 30 predicted N-linked glycosylation sequences (Asn-X-Thr or Asn-X-Ser) for the parental strains compared to 27 (NC7) and 29 (JC16) for the progeny viruses. Most of the glycosylation sites reside in the SU portion of Env with HIV-1_{SF2} containing the highest number and HIV-1 isolate NC the least (25 for SF2, 24 for LAI and JC, and 23 for NC). Glycosylation of glycoproteins has been shown to influence the immune response toward virus infection. As expected, the gp120 (SU) glycoprotein contained the highest number of mutations. The V1-V2, V3, V4, and V5 hypervariable regions contained 10, 8, 8, and 3 point mutations respectively specific to JC16 and NC7. The V1-V2 region of JC16 and NC7 also contained insertions relative to the other viruses, resulting in amino acid lengths of 72 (progeny viruses), 69 (LAV), 70 (SF2), and 61 (NDK). There were multiple amino acid deletions in the V4 regions and single amino acid insertions in V5 regions of JC16 and NC7 relative to the parental strains. Both the CD4binding domain and the proteolytic cleavage site (REKR) at the SU/TM junction were perfectly conserved.

The V3 region was the most interesting of the hypervariable regions. While the parental strains contained only 9 basic amino acids (Arg, Lys, and His positively charged residues) JC16 and NC7 HIV-1 isolates had 12 basic and 2 negatively charged residues (Asp and Glu). This gave the progeny viruses a net positive charge of 10 in the entire V3 region and an overall positive charge of +1 (LAV) or +2 (SF2) compared to the parental

strains. Eight of the 10 resultant positive charges for JC and NC isolates are located between residues 10 and 27 of V3 (Fig. 3) compared with 5 of 9 (LAV) and 4 of 9 (SF2) in the same region. At least within this region, JC and NC isolates seem to have a relatively high net positive charge of 4 (relative to SF2) and 3 (relative to LAV). Other researchers have shown that changes in basic amino acids in the middle portion of V3 loop (residues 10 to 27 in Table 5) can alter the syncytium-inducing properties and phenotype of the virus [Bhattacharyya et al. (1996) AIDS Res. Hum. Retroviruses 12, 83-90; De Wolf et al. (1994) AIDS Res. Hum. Retroviruses 10, 1387-1400; Okada et al. (1994) AIDS Res. Hum. Retroviruses 10, 803-811].

To study the biological activities of the cloned and uncloned viruses derived from C499 and C455, we conducted in vitro replication studies. Figures 4A, 4B and 4C shows the results of replication studies in mitogen-stimulated cPBMC. Two of the three viruses used for inoculation of C499, SF2 and LAV-1b, were able to replicate in stimulated cPBMC, albeit with different kinetics (Fig. 4A). The SF2 isolate grew very slowly and to low titers in cPBMC. In contrast, the LAV-1b isolate grew very well and with rapid kinetics, with a high titer of virus already present by day 7 post infection. The SF2 isolate was unable to induce detectable syncytium formation in cPBMC. The LAV-1b isolate, under these conditions, induced very few syncytia, in contrast to previously reported results [Watanabe et al. (1991) J. Virol. 65, 3344-3348]. Included in these analyses was the DH12 isolate of HIV-1 [Shibata et al. (1995) J. Virol. 69, 4453-4462]. This primary isolate from a human has been shown to be highly cytopathic for cPBMC. While this virus quickly established infection in the stimulated cell population, it did not grow to high titers. The numerous syncytia formed infection with DH12 could account for the lack of growth observed. The uncloned and cloned viruses of JC (Fig. 4B) and NC (Fig. 4C) replicated to levels comparable to those of LAV-1b for the same period of time. The rates of replication for cloned and uncloned viruses were indistinguishable.

The ability of HIV-1 virus isolates to replicate in unstimulated cPBMC was similarly evaluated. Results of this assay (Fig. 4D, 4E and 4F) showed that only the NC (cloned and uncloned) and the LAV-1b isolates of HIV-1 were capable of significant replication in unstimulated cPBMC. Replication rates for these viruses were less than 10% those observed

in stimulated cPBMC. Additionally, the kinetics of virus production in unstimulated cells was much slower than that observed in stimulated cPBMC. Interestingly, JC (cloned and uncloned) isolates of HIV-1 failed to replicate in cPBMC, reflecting an inherent biological difference between the JC and NC viruses. While the DH12 isolate was able to replicate in unstimulated cells, the levels of virus achieved were much less than that of the other viruses. The SF2 isolate was unable to replicate in unstimulated PBMC. Virus recovered from the molecular clones displayed the intrinsic replicative properties exhibited by the viruses they were derived from.

Reports of HIV-1 isolates able to replicate in chimpanzee macrophages has been controversial. To investigate the ability of these viruses and their respective clones to replicate in macrophages, we conducted in vitro assays using purified chimpanzee monocytederived macrophages (MDM). Figure 5 shows virus production in macrophages at 7 and 14 days post infection. Both the HIV-1 JC and NC virus isolates (cloned and uncloned) replicated in MDM as determined by the levels of p24 antigen produced. However, the amount of virus produced by the HIV-1_{JC16} molecular clone at 14 days post infection was twice that produced by the HIV-1_{JC}. The titers for cloned and uncloned HIV-1_{NC} viruses were comparable for the same period of time. Among the viruses that were tropic for MDM, HIV-1_{IC} (uncloned) produced the least amount of virus. HIV-1_{LAV-1b} and HIV-1_{DH12} also infected MDM and produced virus which is consistent with previous observations made by Gendelman et al. [Gendelman et al. (1991) *J. Virol.* 65, 3853-3863] and Shibata et al. [Shibata et al. (1995) *J. Virol.* 69, 4453-4462] respectively. Unlike HIV-1_{LAV-1b}, the other parental virus, HIV-1_{SF2}, did not replicate in MDM.

Because the cloned viruses displayed such similar in vitro biologic activities as the uncloned stocks, we sought to examine whether these viruses represented major species in the viral mix. Genomic DNA was isolated and nested PCR was used to amplify the V1-V2 and V3-V5 regions of HIV-1 envelope gene. These hypervariable regions of env gene were selected because most viral heterogeneity has been associated with them. Equal amounts of PCR products were mixed, heat denatured, and then reannealed with analogous fragments derived from standards (SF162 subtype B3 and ZM18 subtype C2) or from JC16 and NC7

molecular clones. Electrophoresis of these fragments on polyacrylamide non-denaturing gels revealed differences in migration rates with homoduplexes moving faster in the gel than heteroduplexes. Homoduplexes based on genomic DNA (HIV-1_{JC} or HIV-1_{NC}) resulted in homoduplex (lower) and heteroduplex (upper) bands that had comparable intensities and were close together lane. The HIV-1_{JC}/HIV-1_{JC16} (but not HIV-1_{NC}/HIV-1_{NC7}) heteroduplex band was between the homoduplex and heteroduplex bands obtained with genomic DNA homoduplexes. The distance between homoduplex and heteroduplex bands formed between JC16 and NC7 molecular clones was larger than that between bands formed from homoduplexes derived from genomic DNA. The SF162 (subtype B3) heteroduplex bands migrated at almost the same rate as the ssDNA but the ZM18 (subtype C2) heteroduplex bands had the slowest migration.

An advantage of establishing a model system for AIDS in smaller nonhuman primates is the decreased cost associated with the housing and upkeep of smaller animals. Animal AIDS models include, but are not limited to, chimpanzees, the gibbons, pig-tailed macaques, and rhesus macaques infected with the HIV-1_{IC} or HIV-1_{NC} virus of the invention.

Disclosed herein are an immunodeficient pig-tailed macaque and also an immunodeficient rhesus macaque; both infected with the HIV-1_{IC} virus of the invention isolated from the C499 chimpanzee which exhibited HIV-1-induced AIDS or the HIV-1_{NC} virus isolated from the C455 chimpanzee. The macaques are infected with HIV-1 isolate JC or HIV-1_{NC} and are used as an animal model for human AIDS in essentially the same manner as chimpanzees as described above. The transmission of HIV-1_{IC} or the HIV-1_{NC} virus into a macaque proceeds via injection of an isolated viral suspension or transfection of a biological fluid or tissue specimen from an HIV-1-infected and AIDS-bearing primate. The HIV-1-infected donor specimen is introduced into a recipient by any suitable means, such as intraperitoneal injection, intravenous injection, surgical implantation and combinations thereof. Donor tissue may be introduced as organized tissue (e.g., thymus, lymph node, etc.) or as discrete cells.

Antigenic fragments of the present invention are peptides which contain at least one epitope (antibody binding site) which binds antibodies which bind to at least one HIV-1 isolate of the present invention. The antigenic fragments are preferably capable of inducing an immune response when administered to a nonhuman primate. DNA encoding such antigenic fragments may be used to transform host cells to thereby produce such antigenic fragments.

Antigenic fragments may be identified by a variety of means. A protein from HIV-1_{JC} and/or from HIV-1_{NC}, such as an envelope protein, may be fragmented with a protease, and the fragments tested to determine whether or not various ones react with antiserum against the protein. See, e.g., J. Robinson et al., Mol. Cell Biochem. 21:23-32 (1978). Another technique is to synthesize peptides which are fragments of the entire protein and determine whether the individual fragments are recognized by neutralizing antibodies against the protein. See, e.g., J. Gerin et al., in Vaccines 85: Molecular and chemical Basis of Resistance to Parasitic, Bacterial and Viral Diseases, 235-239 (Lerner et al., eds. 1985). Still another method useful for obtaining immunogenic fragments of a protein is by isolation and identification of monoclonal escape mutants. In this strategy, HIV-1 is produced in the presence of a monoclonal antibody to the virus. The only viruses which can grow under these conditions are those with a mutation in the nucleotide sequence which codes for an epitope to which the monoclonal antibody binds. A mutant virus which grows under these conditions is referred to as the "monoclonal escape mutant." The monoclonal escape mutant is then sequenced and the mutant sequence compared with the nucleotide sequence of the HIV-1_{3C} isolate or the HIV-1_{NC} isolate to find the specific location of the mutation. The mutation is located in a region which codes for a protective epitope, or an "immunogenic fragment." See, e.g., J. Lopez et al., J. Virol. 64:927 (1990).

Antigenic preparations of the present invention are useful as reagents in immunoassay diagnostic studies of retroviruses. Immunochemical methods for detecting retroviruses include, for example, immunofluorescence assays or immunoenzymatic assays. Immunofluorescence assays typically involve incubating, for example, serum from the subject to be tested with preparations of the pathogenic virus or fragments thereof. Immune

complexes formed are detected using either direct or indirect methods, for example, the use of antibodies to which fluorescent labels such as rhodamine or fluorescein have been coupled. Immunoenzymatic assays typically involve viral extracts or other antigen-containing compositions bound to a surface. Serum from a subject to be tested for the presence of antibodies directed against one or more antigens is contacted with the surface and, after a period of incubation, unbound substances are washed away. The presence of immune complexes is detected using antibodies labeled with an enzyme such as horseradish peroxidase, alkaline phosphatase, or beta-galactosidase, which is capable of converting a colorless or nearly colorless substrate into a highly colored product, or an enzyme which emits light in the presence of the proper substrate. The amount of product formed is detected visually, spectrophotometrically, or luminometrically and is compared to a similarly treated control. The presence of antibodies in biological fluids may also be detected by agglutination. Viral lysates or antigen compositions are used to coat, for example, latex particles.

Diagnostic tests utilizing the present invention may be carried out in accordance with known techniques. Such techniques provide a method of detecting the presence of HIV-1 by detecting the presence of HIV-1 antibodies. Such methods comprise collecting an antibody-containing biological sample (e.g., blood, blood sera, blood plasma, cerebrospinal fluid, tissue samples) from the subject, contacting the sample with an antigenic preparation of the viral particles of the present invention as given herein, and then detecting the formation of a reaction product between the antibodies in the sample and the antigenic preparation. Any suitable assay format, such as radioimmunoassay (RIA) or enzyme-linked immunosorbent assay (ELISA) may be employed, in accordance with known techniques. See, e.g., Immunology: Basic Processes, 162-175 (J. Bellanti, [Ed.] 2d ed., W.B. Saunders Co. (1985).

Also disclosed herein are kits for the detection of HIV infection. Such kits comprise a container containing an antigenic preparation of the viral particles of the present invention, which may be lyophilized. The antigenic preparation may comprise, e.g., the HIV-1 envelope protein (env) or the group antigen (gag) protein of the HIV-1_{IC} and/or HIV-1_{NC} of the invention.

A method of inducing antibodies to HIV-1 in a subject, as disclosed herein, comprises administering to a subject an immunogenic amount of infectious viral particles of the present invention. This method may be used to make polyclonal or monoclonal antibodies, which may be used in diagnostic assays. Suitable subjects include mammals (such as, for example, rats, rabbits, mice, and horses) and primates. The term primates is herein intended to encompass any members of the order Primata (for example, lemurs, mandrills, rhesus monkeys, macaques, and chimpanzees) and to include humans. Suitable subjects include those in which antibodies to HIV may be raised (e.g., rabbit, horse).

In the above methods of inducing antibodies, viral antigenic preparations of the present invention may be combined with any suitable pharmaceutically acceptable carrier (such as sterile, pyrogen-free physiological saline solution, or sterile, pyrogen-free phosphate-buffered saline solution). The viral antigens are included in an effective immunogenic amount. The precise amount to be administered to a given subject is determined by techniques known in the art and will vary depending on the route of administration, the subject and the desired response. Administration to the subject may occur by any suitable route (e.g., by intramuscular injection, subcutaneous injection, intraperitoneal injection, or intravenous injection). The appropriate immunogenic dosage will depend upon the particular subject and the desired outcome. Techniques to determine a particular immunogenic amount of the viral particles of the present invention will be apparent to those of ordinary skill in the art. See, e.g., Johnson et al., *Proc. Natl. Acad. Sci. USA* 89:2175 (1992). For example, the active agent (viral particles or preparations thereof) may be given in an amount of from 0.05 to 50 µg per kg body weight (e.g., 0.5 or 1.0 µg per kg).

The invention also provides for a variety of different vaccines based on the structures of the HIV-1_{JC} isolate and/or the HIV-1_{NC} of the invention and a method for vaccinating a population against HIV. Examples of active agents used for the preparation of a vaccine of the invention include the live attenuated HIV-1_{JC} and/or HIV-1_{NC} isolate, fixed whole virus, host cells expressing virus antigen, preparations of virus fragments, purified proteins, antigenic fragments of proteins and antigenic peptides which are derivatives of the antigenic fragments. According to the present invention, HIV-1_{JC}- and/or HIV-1_{NC}-derived

compositions or vaccines are useful for preinfection immunization of primates as well as for postinfection (therapeutic) immunization of HIV-infected primates (see Fultz et al. [1989], Lerner et al., eds. Cold Spring Harbor, NY).

Live attenuated HIV isolate JC virus (or HIV-1_{NC} virus) is prepared by serial passage of the virus in tissue culture or genetically altered by recombinant techniques, in accordance with known procedures. Fixed virus is made by contacting live virus (attenuated or unattenuated) to a suitable fixative, such as formalin.

Preparations of viral fragments are made by lysing host cells, such as *E. coli* cells, transformed with a vector encoding an HIV-1 isolate of the present invention (or both) or a portion thereof. The lysate may be used in crude or partially purified form, or a particular viral protein (or antigenic fragment thereof) such as the envelope protein, can be purified to homogeneity and used as an active agent for a vaccine against HIV-1. Host cells such as yeast cells may be transformed with vectors of the present invention capable of expressing HIV-1 proteins, or antigenic fragments thereof, on the surface of the host cells, and the transformed host cells used as an active vaccine agent as such or fixed (e.g., with formalin) and used as an active agent.

Antigenic peptides are selected from the group consisting of antigenic fragments of HIV isolate JC and/or NC proteins, such as the envelope protein, and the antigenic equivalents thereof (i.e., analogs or derivatives). Antigenic peptides may be chemically synthesized or produced by recombinant techniques.

Viral antigenic preparations and cells producing viral antigens and/or fragments thereof may be formulated into immunogenic compositions as neutral or salt forms. Preferably, when cells are used they are of avirulent strains, or the cells are killed before use. Pharmaceutically acceptable salts include but are not limited to the acid addition salts (formed with free amino groups of the peptide) which are formed with inorganic acids, e.g., hydrochloric acid or phosphoric acids; and organic acids, e.g., acetic, oxalic, tartaric, or maleic acid. Salts formed with the free carboxyl groups may also be derived from inorganic

bases, e.g., sodium, potassium, ammonium, calcium, or ferric hydroxides, and organic bases, e.g., isopropylamine, trimethylamine, 2-ethylamino-ethanol, histidine, and procaine.

The term "antigenic equivalents," as used herein, refers to proteins or peptides which bind to an antibody which binds to the protein or peptide with which equivalency is sought to be established. Antibodies which are used to select such antigenic equivalents are referred to as "selection antibodies" herein. Preferred selection antibodies are monoclonal antibodies which bind to HIV isolate JC and/or to HIV-1_{NC}, but not to prior isolates of HIV-1 such as the HIV-1 isolates NDK, LAI and SF2, for example.

One or more amino acids of an antigenic peptide sequence may be replaced by one or more other amino acids which do not affect the antigenicity of that sequence. Such changes can be guided by known similarities between amino acids in physical features such as charge density, hydrophobicity/hydrophilicity, size and configuration, For example, threonine and serine can be interchanged, or aspartic acid and glutamic acid, or leucine and isoleucine, and the like.

Antigenic equivalents may be formed by modifying reactive groups within a natural sequence or modifying the N-terminal amino and/or C-terminal carboxyl group. Such equivalents include salts formed with acids and/or bases, particularly physiologically acceptable inorganic and organic acids and bases. Other equivalents include modified carboxyl and/or amino groups on the synthetic peptide to produce esters or amides, or amino acid protecting groups such as N-t-butoxycarbonyl. Preferred modifications are those which provide a more stable, active peptide which will be less prone to enzymatic degradation in vivo.

In a particular embodiment of the invention, polyclonal and/or monoclonal antibodies capable of specifically binding to a particular epitope of at least one HIV-1 isolate of the invention are provided. The term *antibody* is used to refer both to a homogenous molecular entity (monoclonal antibody) and a mixture (such as a serum product) made up of a plurality of different molecular entities (polyclonal antibody). Monoclonal or polyclonal antibodies,

and preferably monoclonal, specifically reacting with a particular epitope of interest can be made by methods known in the art. See, e.g., Harlow and Lane (1988) *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratories; Goding (1986) *Monoclonal Antibodies: Principles and Practice*, 2d ed., Academic Press, New York. Also, recombinant immunoglobulins may be produced by methods known in the art, including but not limited to the methods described in U.S. Patent No. 4,816,567, incorporated by reference herein.

Monoclonal antibodies with affinities of 10⁸ M⁻¹, preferably 10⁹ to 10¹⁰ or more are preferred.

For use as a vaccine, immunogenic compositions may be formulated by any of the means known in the art. Such vaccines are typically prepared as injectables, either as liquid solutions or suspensions. Solid forms suitable for solution in, or suspension in, liquid prior to injection may also be prepared. The preparation may also, for example, be emulsified, or the protein encapsulated in liposomes. Such vaccines may be administered to the subject by any suitable means, for example, by intramuscular injection, by subcutaneous injection, by intraperitoneal injection, by oral injection, and by nasal spray.

The vaccine or other immunogenic composition may be given in a single dose or multiple dose schedule. A multiple dose schedule is one in which a primary course of vaccination may include 1 to 10 or more separate doses, followed by other doses administered at subsequent time intervals as required to maintain and or reinforce the immune response, e.g., at 1 to 4 months for a second dose and, if needed, a subsequent dose(s) after several months.

The immunogenic peptide antigen compositions are administered in a manner compatible with the dosage formulation and in such amount as will be prophylactically and/or therapeutically effective. The quantity to be administered, which is generally in the range of about 100 to 1,000 μ g of protein per dose, more generally in the range of about 5 to 500 μ g of protein per dose, depends on the subject to be treated, the capacity of the individual's immune system to synthesize antibodies, and the degree of protection desired. Precise amounts of the active ingredient required to be administered may depend on the judgment of the physician

and may be peculiar to each individual, but such a determination is within the skill of such a practitioner.

Vaccine formulations of the present invention comprise the active agent mixed with excipients or carriers which are pharmaceutically acceptable and compatible with the active ingredient. Suitable excipients include but are not limited to water, saline, dextrose, glycerol, ethanol, or the like and combinations thereof. The concentration of the immunogenic polypeptide in injectable formulations is usually in the range of 0.2 to 5 mg/ml.

In addition, if desired, the vaccines may contain minor amounts of auxiliary substances such as wetting or emulsifying agents, pH buffering agents, and/or adjuvants which enhance the effectiveness of the vaccine. Examples of adjuvants which may be effective include but are not limited to: aluminum hydroxide; aluminum phosphate; plant and animal oils; synthetic polymers; e.g., N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP); N-acetyl-nor-muramyl-L-alanyl-D-isoglutamine (CGP 11637, referred to as nor-MDP); N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphosphoryloxy)-ethylamine (CGP 19835A, referred to as MTP-PE); and RIBI, which contains three components extracted from bacteria, monophosphoryl lipid A, trehalose dimycolate; cell wall skeleton (MPL+TDM+CWS) in a 2% squaline/Tween 80 emulsion; etc.

In addition, the vaccine formulations may also contain one or more stabilizer, for example, carbohydrates such as sorbitol, mannitol, starch, sucrose, dextrin, and glucose, proteins such as albumin or casein, and buffers such as alkaline metal phosphates and the like.

In view of the similarities in protein molecules making up different isolates of HIV-1, the skilled artisan understands that an antibody, particularly a monoclonal antibody, which is specific for a particular epitope directed to a particular protein of the HIV-1_{IC} isolate and/or the HIV-1_{NC} isolate, can be used to screen for other HIV-1 isolates having similar epitopes recognized by that (monoclonal) antibody.

Antibodies generated against specific epitopes of the HIV-1_K of the invention are useful, for example, as probes for screening DNA expression libraries or for detecting the presence of HIV-1 strains in a test sample. Antigens can be synthesized and conjugated to a suitable carrier protein (e.g., bovine serum albumin or keyhole limpet hemocyanin) for use in vaccines or in raising specific antibodies. Frequently, the polypeptides and antibodies will be labeled by joining, either covalently or noncovalently, a substance which provides a detectable signal. Suitable labels include but are not limited to radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent agents, chemiluminescent agents, magnetic particles and the like. United States Patents describing the use of such labels include but are not limited to Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,355,241.

Immunogenic carriers may be used to enhance the immunogenicity of an active agent. Such carriers include but are not limited to proteins and polysaccharides, liposomes, and bacterial cells and membranes. Protein carriers may be joined to the capsular polysaccharide molecules to form fusion proteins by recombinant or synthetic means or by chemical coupling. Useful carriers and means of coupling such carriers to polypeptide antigens are known in the art. The art knows how to administer immunogenic compositions so as to generate protective immunity where immunity is most helpful.

Compositions and immunogenic preparations including vaccine compositions comprising substantially purified antigens derived from an HIV-1 isolate JC and/or NC and a suitable carrier therefor are provided. Immunogenic compositions are those which result in specific antibody production when injected into a human or an animal. Such immunogenic compositions are useful, for example, in immunizing primates against infection by HIV-1 strains. The immunogenic preparations comprise an immunogenic amount of, as specifically exemplified, at least one antigenic determinant derived from the HIV-1_{IC} isolate of the invention and a suitable carrier. Alternatively, the immunogenic composition can comprise host cells harboring an antigenic agent from the specifically exemplified HIV-1_{IC} strain and a suitable carrier. It is understood by one of ordinary skill in the art that a functionally equivalent, recombinant mutant of HIV-1_{IC} and or HIV-1_{NC} can be produced by the

introduction of the cloned DNA containing the insertion mutations responsible for a desired characteristic. It is also within the scope of the present invention and readily within the grasp of the ordinary skilled artisan to generate other types of genetically stable mutations in the structural or enzyme genes of HIV-1. Such immunogenic compositions (or vaccines) are useful, for example, in immunizing an animal, especially humans, against AIDS and related diseases resulting from infection by HIV-1 species. Such immunogenic compositions can also elicit the production of antibodies which will cross react with proteins of other HIV-1 and HIV-2 strains expressing epitopes in common with those of the starting HIV-1_{JC} isolate. It is understood that where whole cells are formulated into the immunogenic composition, the cells are preferably inactivated, especially if the cells are of a virulent strain. Such immunogenic compositions may comprise one or more protein or the immunogenic cellular component. By "immunogenic amount" is meant an amount capable of eliciting the production of antibodies directed against an antigenic agent of HIV-1_{JC} in an animal or human to which the vaccine or immunogenic composition has been administered.

The nucleotide sequence of the HIV-1_{JC} isolate or the HIV-1_{NC} isolate can be used to generate hybridization probes which specifically bind to HIV-1_{JC} genetic material, or to DNA of HIV-1 isolates having the identifying characteristics of the HIV-1 isolates JC or NC, to determine the presence of such HIV-1 in primates. The hybridization probe may be selected so that it does not bind to other known HIV-1 isolates such as NDK, LAI, SF2, etc. The hybridization probes can be cDNA fragments or polynucleotides and may be labeled with a detectable group, as is well-known in the art. Pairs of probes can serve as PCR primers for synthesis and amplification processes in accordance with the description, for example in U.S. Patent Nos. 4,683,202 and 4,683,195.

In specific embodiments, probes of the invention comprise DNA sequences of HIV- $l_{\rm IC}$ or sequences encoding antigenic fragments thereof or sequences having identity thereto. In particular are provided probes having a DNA sequence as set forth in SEQ ID NO:1 or a sequence having identity thereto.

The production of DNA, vectors, transformed host cells, HIV-1 virus, proteins, and protein fragments of the present invention by genetic engineering techniques can be carried out in accordance with methods known in the art. See, e.g., U.S. Pat. No. 4,761,371, U.S. Pat. No. 4,877,729, U.S. Pat. No. 4,912,038, and U.S. Pat. No. 4,879,224, among others.

A nucleotide sequence (polynucleotide) or fragment thereof is substantially homologous (or substantially similar) to another polynucleotide if, when optimally aligned (with appropriate nucleotide insertions or deletions) with another polynucleotide, there is nucleotide sequence identity for approximately 80% of the nucleotide bases, usually approximately 90%, more preferably about 95% to 100% of the nucleotide bases. Gaps introduced to optimize alignments are treated as mismatches.

Alternatively, substantial homology (or similarity) exists when a polynucleotide or fragment thereof will hybridize to another polynucleotide under selective or stringent hybridization conditions. Selectivity of hybridization exists under stringent hybridization conditions which allow one to distinguish the target polynucleotide of interest from other polynucleotides. Typically, selective hybridization will occur when there is approximately 75% similarity over a stretch of about 14 nucleotides, preferably approximately 80% similarity, more preferably approximately 85% similarity, and most preferably approximately 90% similarity. See Kanehisa (1984) *Nucl. Acids Res.*, 12:203-213. The length of homology comparison, as described, may be over longer stretches, and in certain embodiments will often be over a stretch of about 17 to 20 nucleotides, preferably 21 to 25 nucleotides, more preferably 26 to 35 nucleotides, and more preferably about 36 or more nucleotides.

The hybridization of polynucleotides is affected by such conditions as salt concentration, temperature, or organic solvents, in addition to the base composition, length of the complementary strands, and the number of nucleotide base mismatches between the hybridizing polynucleotides, as will be readily appreciated by those skilled in the art. Stringent temperature conditions will generally include temperatures in excess of 30°C, typically in excess of 37°C, and preferably in excess of 45°C. Stringent salt conditions will ordinarily be less than 1 M, typically less than 500 mM, and preferably less than 200 mM.

However, the combination of parameters is much more important than the measure of any single parameter (Wetmur and Davidson [1968] J. Mol. Biol. 31:349-370).

An isolated or substantially pure polynucleotide is a polynucleotide which is substantially separated from other polynucleotide sequences which naturally accompany a native sequence. The term embraces a polynucleotide sequence which has been removed from its naturally occurring environment, and includes recombinant or cloned DNA isolates, chemically synthesized analogues and analogues biologically synthesized by heterologous systems.

A polynucleotide is said to encode a polypeptide if, in its native state or when manipulated by methods known to those skilled in the art, it can be transcribed and/or translated to produce the polypeptide of a fragment thereof. The anti-sense strand of such a polynucleotide is also said to encode the sequence.

Vectors are replicable DNA constructs used to either amplify or express DNA of the present invention. An expression vector is a replicable DNA construct in which DNA of the present invention is operably linked to control sequences capable of expressing that DNA in a suitable host. Generally, control sequences include a transcriptional promoter, an optional operator sequence to control transcription, a sequence encoding suitable mRNA ribosomal binding sites, and sequences which control the termination of transcription and translation. Suitable vectors include plasmids, viruses (e.g., vaccinia virus, adenovirus, baculovirus, cytomegalovirus) phage, and integratable DNA fragments (i.e., fragments integratable into the host genome by recombination).

DNA regions are operably linked or operably associated when they are functionally related to each other. For example, a promoter is operably linked to a coding sequence if it controls the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to permit translation.

Transformed host cells are cells which have been transformed or transfected with vectors as described above. Transformed host cells ordinarily express the DNA of the present invention. Suitable host cells include prokaryote, yeast or higher eukaryotic cells such as mammalian cells and insect cells.

Prokaryote host cells include gram negative or gram positive organisms, for example Escherichia coli (E. coli) or Bacilli. Exemplary host cells are E. coli W3110 (ATCC 27,325), E. coli B, E. coli X1776 (ATCC 31,537), E. coli 294 (ATCC 31,446). A broad variety of suitable prokaryotic and microbial vectors are available. E. coli is typically transformed using pBR322. Promoters most commonly used in recombinant microbial expression vectors include the β-lactamase (penicillinase) and lactose promoter systems (Change et al., Nature 275:615 [1978]; and Goeddel et al., Nature 281:544 [1979], a tryptophan (trp) promoter system (Goeddel et al. [1980] Nuc. Acids Res. 8:4057, and EPO App. Publ. No. 36,766) and the tac promoter (H. De Boer et al., Proc. Natl. Acad. Sci. USA 80:21 [1983]). The promoter and Shine-Dalgarno sequence are operably linked to the DNA of the invention, i.e., they are positioned so as to promote transcription of messenger RNA from the DNA.

Eukaryotic microbes such as yeast cultures may also be transformed with vectors of the present invention. See, e.g., U.S. Pat. No. 4,745,057. Saccharomyces cerevisiae is the most commonly used yeast, although other yeast may also be used. Host cells such as insect cells (e.g., cultured Spodoptera frugiperda cells) and expression vectors such as the baculovirus expression vector may be employed in carrying out the present invention, as described in U.S. Pat. Nos. 4,745,051 and 4,879,236 to Smith et al.

Examples of useful mammalian host cell lines are VERO and HeLa cells, Chinese hamster ovary (CHO) cell lines, and WI138, BHK, COS-7, CV, and MDCK cell lines. The transcriptional and translational control sequences in expression vectors to be used in transforming vertebrate cells are often provided by viral sources. For example, commonly used promoters are derived from polyoma, Adenovirus 2, and Simian Virus 40 (SV40). See, e.g., U.S. Pat. No. 4,599,308. An origin of replication may be provided either by construction of the vector to include an exogenous origin, such as may be derived from SV40 or other viral

(e.g., Polyoma, Adenovirus, VSV, or BPV) source, or may be provided by the host cell chromosomal replication mechanism. If the vector is integrated into the host cell chromosome, the latter is often sufficient. Rather than using vectors which contain viral origins of replication, one can transform mammalian cells by the method of cotransformation with a selectable marker and DNA of the present invention, as described in U.S. Pat. No. 4,399,216.

Except as noted hereafter, standard techniques for cloning, DNA isolation. amplification and purification, for enzymatic reactions involving DNA ligase, DNA polymerase, restriction endonucleases and the like, and various separation techniques are those known and commonly employed by those skilled in the art. A number of standard techniques are described in Sambrook et al. (1989) Molecular Cloning, Second Edition, Cold Spring Harbor Laboratory, Plainview New York; Maniatis et al. (1982) Molecular Cloning, Cold Spring Harbor Laboratory, Plainview, New York; Wu (ed.) (1993) Meth. Enzymol. 218:Part I; Wu (ed.) (1979) Meth. Enzymol. 68; Wu et al. (eds.) (1983) Meth. Enzymol. 100 and 101; Grossman and Moldave (eds.) Meth. Enzymol. 65; Miller (ed.) (1972) Experiments in Molecular Genetics, Cold Spring Harbor Laboratory, Plainview, New York; Old and Primrose (1981) Principles of Gene Manipulation, University of California Press, Berkely; Schleif and Wensink (1982) Practical Methods in Molecular Biology; Glover (ed.) (1985) DNA Cloning Vol. I and II, IRL Press, Oxford, UK; Hames and Higgins (eds.) (1985) Nucleic Acid Hybridization, IRL Press, Oxford, UK; and Setlow and Hollaender (1979) Genetic Engineering: Principles and Methods, Vols. 1-4, Plenum Press, New York. Abbreviations and nomenclature, where employed, are deemed standard in the field and commonly used in professional journals such as those cited herein.

All publications, patent applications and patents cited herein are incorporated by reference to the extent that they are not inconsistent with the present disclosure.

The foregoing discussion and the following examples are provided for illustrative purposes, and they are not intended to limit the scope of the invention as claimed herein.

Modifications and variations which may occur to one of ordinary skill in the art are within the

intended scope of this invention. The present invention is further described in the nonlimiting examples set forth below.

EXAMPLES

Example 1. Animal Subjects

All animals (e.g., chimpanzees) were maintained in accordance with the guidelines established by the Animal Welfare Act and the NIH guide for care and use of laboratory animals. The Yerkes Center is fully accredited by the American Association for the Accreditation of Laboratory Animal Care (AAALAC).

A cohort of 12 chimpanzees was inoculated with several strains of HIV-1 at the Yerkes Center. A member of this cohort, chimpanzee 499 was inoculated on three different occasions with three different HIV-1 isolates: HIV-1_{SF2} in 1985, HIV-1_{LAI} in 1986 and HIV-1_{NDK} in 1987 (See Fultz et al. [1991] J. Infect. Dis. 163:441-447 and Novembre et al. [1997] J. Virol. 71:4086-4091).

It is preferred that a chimpanzee be bred in captivity, be seropositive for Epstein-Barr virus and cytomegalovirus and be seronegative and PCR negative for HIV prior to being used as an animal model.

Example 2. <u>Isolation and Purification of HIV-1_{IC} Isolate</u>

The original source of the HIV-1_{JC} isolate was the chimpanzee C499. HIV was easily isolated from the peripheral blood mononuclear cells (PBMC) of this animal. Cocultivation of PBMC derived from C499 with uninfected chimpanzee PBMC (cPBMC) resulted in the isolation of a virus (HIV-1_{JC}) which induced syncytium formation in chimpanzee cells. The nucleotide sequence of this virus was distinguished from the isolates used for the initial inoculations and from other known HIV-1 isolates.

The HIV-1_{IC} isolate was deposited with the AIDS Reagent Program, McKesson Bioservices, 685 Lofstrand Lane, Rockville, MD 20850 USA, a division of the NIH AIDS

Research and Reference Reagent Program, in May 1997 and has been assigned Catalog Number 3523. A substantially full-length infectious molecular clone has a nucleotide sequence as given in SEQ ID NO:11.

In a specific embodiment of the invention, probes of the invention comprise DNA sequences as set forth in SEQ ID NO:11 and/or sequences of at least 15 contiguous nucleotides derived therefrom or sequences complementary thereto.

The original source of the HIV-1_{JC} isolate was the chimpanzee C455. HIV was easily isolated from the peripheral blood mononuclear cells (PBMC) of this animal. Cocultivation of PBMC derived from C455 with uninfected chimpanzee PBMC (cPBMC) resulted in the isolation of a virus (HIV-1_{NC}) which induced syncytium formation in chimpanzee cells. The nucleotide sequence of this virus is distinguished from other known HIV-1 isolates. A substantially full-length infectious molecular clone has a nucleotide sequence as given in SEQ ID NO:12.

In a specific embodiment of the invention, probes of the invention comprise DNA sequences as set forth in SEQ ID NO:12 and/or sequences, of at least 15 contiguous nucleotides derived therefrom or sequences complementary thereto.

Example 3. Virus and Viral Antigens

The HIV-1 isolates used to inoculate C499 included LAV (lymphadenopathy-associated virus)-1_{BRU} (Barre-Sinoussi [1983] Science 220:868-871), SF2 (previously designated ARV [AIDS-related virus]-2) (Levy et al. [1984] Science 225:840-842), and NDK, a highly cytopathic HIV-1 of African origin (Spire et al. [1989] Gene 81:275-284). Inoculations of virus were done intravenously with 1-ml aliquots of undiluted or diluted virus stocks. In January 1988, 33 months after inoculation HIV-1_{SF2}, C499 was part of a study to assess the effects of therapeutic vaccination on immunity and viral status in HIV-1-infected chimpanzees. C499 was given two intramuscular injections, 4 weeks apart, of 500 μg of recombinant SF2 p53 gag, produced in yeast and formulated with 100 μg of muramyl

tripeptide (Ciba-Geigy, Summit, NJ) in 4% squalene and 0.008% Tween 80 (Fultz et al. [1989] in Vaccines 89, supra).

The HIV-1 isolates used to inoculate C455 was JC, deposited with the AIDS Reagent Program, Catalog Number 3523. Inoculations of virus were done intravenously with 1-ml aliquots of undiluted or diluted virus stock. C534 was given two intramuscular injections, 4 weeks apart, of 500 µg of recombinant SF2 p53 gag, produced in yeast and formulated with 100 µg of muramyl tripeptide (Ciba-Geigy, Summit, NJ) in 4% squalene and 0.008% Tween 80 (Fultz et al. [1989] in Vaccines 89, supra).

Example 4. Serologic Assays

Serum samples were tested by indirect ELISA for antibodies to specific HIV-1 proteins using a series of recombinant antigens. These antigens have been described (Fultz et al. [1989] in *Vaccines* 89, *supra*) and included p25 *gag* and p53 *gag*, produced in yeast; p31 *pol*; and *env2*-3(SF2) and *env2*-3 (IIIB), nonglycosylated polypeptides corresponding to full-length gpl20. Antibody titers to whole-virus preparations were determined with an HIV enzyme immunoassay kit (EIA; Genetic Systems, Seattle, WA). Neutralization assays were performed as described (Fultz et al. [1986] Proc. Natl. Acad. Sci. 83:5286-5290); titers were based on >80% inhibition of reverse transcriptase (RT) activity after preincubation of virus with serum and infection of normal human PBMC. Antibodies cross-reactive with histone H2B were identified by immunoblot using purified calf thymus histones as described previously (Strickler et al. [1987] Nature 327:710-713).

Example 5. Virus Assays

Assays to detect virus in plasma samples or to recover virus from chimpanzee PBMC were performed with normal human PBMC as indicator cells, unless otherwise indicated, and have been described (Fultz et al. [1986] J. Virol. 58:116-124). Medium was RPMI 1640 with 10% fetal bovine serum, 8 units of recombinant interleukin-2 (IL-2)/ml, glutamine, and antibiotics (RPMI-IL-2). Infectious PBMC were quantified by serial 1:5 limiting dilution and cocultivation with human PBMC (Fultz et al. [1986], *supra*). HIV-1 was detected in all assays by the presence of RT activity in cell-free culture supernatants. To detect HIV antigen in

serum or plasma samples, a commercially available HIV-1 antigen capture kit (Coulter, Hialeah, FL) was used.

To assess the influence of CD8⁺ lymphocytes on viral replication and recovery, after 3 days of stimulation with concanavalin A (ConA 10 μg/ml) in RPMI 1640, PBMC were washed once; 10⁷ cells from C499 were placed in fresh RPMI-IL-2, cultured without indicator cells, and monitored for HIV-1 production. In other experiments, the overnight incubation of C499's PBMC to remove adherent cells, CD4⁺- and CD8⁺-enriched populations were obtained by panning with monoclonal antibodies specific for the CD8⁺ antigen or with CD8⁺-coated magnetic beads (Dynabeads; Robbins Scientific, Mountain View, CA). After removal of the CD8⁻[¹⁷⁷] cells, CD8⁺ lymphocytes were recovered from the plates or magnetic beads by additional overnight incubation and repeated washes with medium. After stimulation with ConA, cultures were established either with CD4⁺-enriched cells only or with CD4⁺- and CD8⁺-enriched cells in various ratios. Percentages of CD4⁺ and CD8⁺ cells in enriched populations were determined by analysis with FACScan (Becton-Dickinson, Mountain View, CA).

Replication kinetics in PBMC and macrophages were tested as follows. 1 x 10⁷ freshly isolated or Con-A stimulated PBMC from HIV-1 negative chimpanzee were infected overnight (at 37°C) with 20 ng of the indicated virus (p24 antigen concentration). The cells were centrifuged at 1000 rpm for 10 min, resuspended in 10 ml complete RPMI medium containing 10% FBS and IL-2 (IL-2 medium) and were incubated at 37°C. Samples of supernatants (1 ml) were harvested on days 3, 7, 10, 14 and 17 post infection. IL-2 medium was added to the cultures following the sampling to maintain the original volume. Supernatants were used in RT assays to determine the relative amounts of virus produced.

For replication in macrophages, cPBMC were resuspended in macrophage media (6 x 10⁶/well in RPMI 1640 containing 15% human serum [AB⁺], 1% HEPES, 0.008 ng/ml GM-CSF, 0.03 ng/ml M-CSF, 1% antibiotic-antimycotic solution (Sigma, St. Louis, MO) and seeded in a 24-well plate and incubated at 37°C for 4 hrs. The cells were mixed by pipetting up and down before the incubation was continued for 4 days. Non-adherent cells were

removed by gently washing the wells. Fresh medium (2 ml/well) was added and the cells were cultured for an additional 3 days to allow full macrophage differentiation. Infections were initiated by adding 10 ng of virus (p24) to the cells in 500 µl media and adsorbed overnight. The inoculum was removed and the cells were washed twice before fresh macrophage medium (2 ml) was added. On days 7 and 14 post infection, aliquots of 0.5 ml were taken for determination of p24 antigen levels using the HIV-1 p24 Antigen kit (Coulter Corp, Miami, FL) according to the manufacturer's instructions.

Example 6. <u>Cell-mediated Immune Responses</u>

Lymphocyte proliferative responses to mitogens were tested by incorporation of [3H]thymidine into the DNA of PBMC seeded in triplicate at 5 x 10⁴ PBMC per well into 96-well plates. PBMC were incubated for four days with different concentrations of phytohemagglutinin (PHA) or ConA, pulsed overnight with 1 μCi of [3H]thymidine per well, and harvested with a cell harvester (Skatron, Sterling, VA); counts per minute (cpm) incorporated were determined in a β counter.

Cytotoxic T lymphocyte (CTL) responses against HIV-1 env- and gag-encoded antigens were assayed using fresh PBMC from C499 as effector cells and autologous Epstein Barr virus-transformed B cells as target cells. Target cells were infected at MOI=10 with recombinant vaccinia viruses expressing HIV-1 gpl 20 env of p24 gag or with wild-type vaccinia as control. Infected targets were labeled with 200 μCi of [51Cr] sodium chromate (Du Pont, Boston) for 2 hours and washed, and 105 target cells (T) were mixed with effector cells (E) at various ratios in 96-well plates. The plates ere incubated for 6 hours at 37°C in 5% CO₂, then 0.1 ml of medium was removed from each well and counted in a γ counter. Maximum release (max) was determined by adding 1% Triton X to labeled target cells, and spontaneous release (spon), by adding only medium. Percentage of specific lysis was determined: [cpm(experimental) - cpm (spon)]/[cpm(max) - cpm(spon)] x 100.

Example 7. Nonhuman Primate Model Systems for AIDS

Nonhuman primates infected with HIV-1 isolate JC or infected with HIV-1_{NC} are useful as a model system for the study of AIDS. Chimpanzees and other monkey species

used for this purpose are preferably specific pathogen-free animals, which are available from primate centers, e.g., the Yerkes Regional Primate Research Center, Emory University, Atlanta, GA.

Infected primates are preferably maintained as a single colony of two or more animals, all inoculated with HIV-1 isolate JC or a colony of two or more animals, all inoculated with HIV-1_{NC}. A colony may be maintained in a single room with each primate housed in an appropriate cage, in accordance with standard practices for the maintenance of animals established by the Animal Welfare Act and the NIH guide for care and use of laboratory animals.

The primates are infected with the HIV-1_{JC} virus or the HIV-1_{NC} virus by any suitable means, such as intraperitoneal, intravenous or subcutaneous injection with a solution containing HIV-1 isolate JC or HIV-1_{NC}. The solution may also be a body fluid or tissue (e.g., blood) from a previously infected primate, a blood fraction containing peripheral blood mononuclear cells from a previously infected primate, a pharmaceutically acceptable carrier such as saline solution containing HIV-1 isolate JC or HIV-1 isolate NC, etc.

Nonhuman primates infected with HIV-_{IC} or HIV-1_{NC} are particularly useful as a model system for AIDS because of the concomitant decrease in CD4⁺ cells and increase in HIV-1 loads in plasma. The development of AIDS has not been previously documented in any nonhuman primate species. When used as a model system, a primate(s) infected with HIV-1_{IC} or HIV-1_{NC} virus is subjected to a treatment useful in combating AIDS in humans and thereafter the progress of the infection and related diseases is monitored. A control (placebo) group of HIV-1_{IC} or HIV-1_{NC}-infected animals is left untreated for comparative purposes. A slowing in the progression of the development of AIDS in infected animals indicates that the treatment may be useful for combating AIDS in humans and additional screening and toxicological testing is prescribed. Such treatment includes but is not limited to a vaccine, a drug (e.g., an antiretroviral compound) or a drug combination (e.g., antiviral nucleosides such as AZT, DDI, etc.), a peptide, a protein, etc.) or a vaccine/drug combination, etc. After treatment initiation, the progress of the disease is monitored by any suitable

parameters including, but not limited to, (a) decline in CD4⁺ cell levels, (b) increase in viral loads in plasma, (c) presence of HIV-1_{IC} virus or HIV-1_{NC} virus, (d) weight loss, (e) general appearance, and other symptoms characteristic of AIDS.

Example 8. Virus Cloning and Sequence Analysis

cPBMC infected with either HIV-1_{JC} or HIV-1_{NC} were used for isolation of DNA using the Puregene Kit (Gentra systems, Minneapolis, MN) as directed by the manufacturer. The strategy used for PCR amplification and primer location on the HIV-1 viral genome is shown in Figure 3. PCR primers MSF12 (5'-AAA TCT CTA GCA GTG GCG CCC GAA CAG-3', (SEQ ID NO:18); HIV-1_{LAV} nt 169 to 195) and MSR5 (5'-GCA CTC AAG GCA AGC TTT ATT GAG GCT-3', (SEQ ID NO:19); HIV-1_{LAV} nt 9225 to 9198) [Salminen et al. (1995) Virology 213, 80-86] were used to amplify a 9056 bp product from PBMC genomic DNA prepared from HIV-1_{sc}-infected cells. Another PCR primer pair 527 (5'-CAC ACA CAA GGC TAC TTC CCT GAT TGG CAG A-3', SEQ ID NO:20, HIV-1_{LAV} nt 5302 to 5274) was used to amplify 5' LTR-containing fragments (5699 bp) from the same DNA source. For the HIV-1_{NC} viral genome, PCR primer pairs 527-528 and 529 (5'-ATG GAA CAA GCC CCA GAA GAC CAA GGG CCA CAG-3', SEQ ID NO:21, HIV-1_{LAV} nt 5141 to 5173) and 530 (5'-GGT CTG AGG GAT CTC TAG TTA CCA GAG TCA C-3', SEQ ID NO:22, HIV-1_{LAV} nt 151 to 121) were used to generate the 5'-half (5699 bp) and 3'-half (4142 bp) PCR products respectively from HIV-1_{NC} genomic PBMC DNA. Primers were synthesized on an Applied Biosystems 392 DNA synthesizer (Applied Biosystems, Foster City, CA). Briefly, PCR was performed using the reagents from the Expand Long Template kit (Boehringer Mannheim, Indianapolis, IN) and 200 ng of DNA template, according to the manufacturer's instructions. After an initial DNA denaturation of 94°C for 2 min, the PCR consisted of 10 cycles of 94°C for 15s, 61°C for 30s, 68°C for 8 min followed by 20 cycles of 94°C for 15s, 61°C for 30s, and 68°C for 8 min with a 5 second addition to each extension. The samples were incubated at 72°C for 30 min after the last cycle and then cooled to 4°C. Results of PCR reactions were evaluated on 0.9% agarose gels. PCR products representing the correct sized fragments were isolated from agarose and were directly cloned into the pCR II plasmid and amplified in Escherichia coli bacteria (TA cloning kit, Invitrogen Corp., San Diego, CA) according to the manufacturer's protocol.

Single bacterial colonies containing plasmids with inserts of the correct size were grown at 30°C overnight and plasmid DNA prepared by the alkaline lysis method.

The strategy for preparing full-length molecular clones is illustrated in Figure 3. Several restriction enzymes were used to generate restriction maps for the positive clones. The Apa I fragment (1947 bp) from the 5' half PCR product was gel-purified and subcloned into the large fragment (7675 bp) of pJC using standard cloning procedures [Sambrook et al. (1989) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY] to generate plasmid pHIV-1_{JC16} (Fig. 1). A chimeric plasmid (pHIV-1_{NCIC}) was generated by subcloning the PCR amplified 5' half of HIV-1_{NC} into pHIV-1_{JC16}. For HIV-1_{NC} full-length clone, the Nco I-Xho I env-containing fragment from the 3' half PCR product was gel purified and subcloned into the Nco I-Xho I large fragment of plasmid pHIV-1_{NCIC} containing the 5' half of HIV-1_{NC}. Multiple restriction enzymes were used for analysis of both viral DNAs to confirm the full length clones.

5 x 106 CEMx174 cells in T-25 flasks were transfected with 2 μg of either pHIV-1_{JC16} or NC7 DNA in transfection buffer (25 mM Tris-HCl)(pH 7.5, 140 mM NaCl, 5 mM Kcl, 0.7 mM K₂HPO₄3H₂O) containing 4 µl of DEAE-dextran (60 mg/ml) for 20 minutes at room temperature. Five milliliters of complete medium (RPMI 1640 supplemented with 10% fetal bovine serum (FBS) and 2 mM L-glutamine) was added to stop the reaction followed by centrifugation of cells at 1000 rpm for 10 minutes. The cells were washed twice in 10 ml complete medium before they were resuspended in 10 ml complete medium, transferred to a T-25 cm² flask and incubated at 37°C (5% CO₂). The cells were checked daily for cytopathic effects (syncytia formation) and aliquots of cultures were tested for the presence of reverse transcriptase (RT) activity using standard assay methods. For 293 cell lines, 2 x 105 cells in 6-well plates were transfected with 2 µg of viral DNA using lipofectin (Life Technologies, Gaithersburg, MD) or DOTAP (Boehringer Mannheim, Indianapolis, IN) according to the manufacturers' instructions. After 24 hours, the transfected cells were overlaid with 2 x 106/well of uninfected cPBMC previously stimulated with concanavalin A (Con-A) for 4 days. After an additional 2 day incubation, the non-adherent cell population (cPBMC) were transferred to a T-25 flask and additional stimulated cPBMC added for virus amplification.

Culture supernatants were assayed for RT activity and the cells were observed daily for development of syncytia. Cell-free stocks of molecularly cloned viruses were prepared at peak RT activity, aliquoted and stored under liquid nitrogen.

Primers for sequencing were constructed from conserved regions of aligned sequences of HIV-1_{LAI} and HIV-1_{SF2} and were synthesized on an applied Biosystems 392 DNA synthesizer. The DNA sequence of each full-length clone virus was determined by the dideoxy-chain termination method using the sequenase system (Amersham Life Sciences, Arlington Heights, IL) and ³⁵S-dATP. Nucleotide sequence alignments were performed with the Intelligenetics Suite of programs (Intelligenetics, Beaverton, OR) whild the phylogenetic analysis of amino acid sequence was done with CLUSTAL method (DNASTAR, Inc., Madison, WI).

Example 9. Heteroduplex Mobility Assays.

The nucleotide sequences for HIV-1_{JC16} and HIV-1_{NC7} have been assigned Genbank accession numbers AF049494 and AF049495 respectively.

The heteroduplex mobility assay kit (NIH AIDS Research and Reference Reagent Program) based on the method described by Delwart et al. (1993) *Science* 262, 1257-1261 was used. Briefly, equal amounts (5 µl each) of second-round PCR products (V1-V2 and V3-V5) from infected cPBMC genomic DNA were mixed with the reference PCR products to obtain heteroduplexes. After adding 1.1 µl of 10x annealing buffer (1 M NaCl, 100 mM Tris [pH 7.8], 20 mM EDTA), the mixed DNAs were denatured at 94 °C for 2 min and then reannealed by rapidly cooling in ice. Three µl of loading dye (25% Ficoll, 1% Orange G) was added to the cooled DNA mixture and the samples loaded onto 5% polyacrylamide gel in 1x TBE (88 mM Tris-borate, 89 mM boric acid, 2 mM EDTA) buffer and electrophoresed at a constant voltage of 250 V for 2.5 hrs. The gels were stained with ethidium bromide and visualized under ultra-violet (UV) light.

TABLE 1 Platelet counts, lymphocyte subset levels, and plasma HIV-1 loads in C499

Date (mo/day/yr)	Platelet count (cells/µl)	Total lymphocyte count (cells µl)*	Absolute CD4* cell count (cells/µl)	Absolute CD8* cell count (cells/µl)	Plasma virus level (RNA equivalents/ml) ^b
10/04/88	63,000	3,136	550	1,020	<104
06/12/89	122,000	3,360	810	1,750	<104
01/02/90	112,000	3,600	1,040	1,690	<104
02/06/90	289,000	5,301_	1,480	2,490	<104
05/11/93	24,000	2,461	390	1,720	0.86 X 10 ⁵
05/18/94	23,000	1,430	300	770	ND⁴
09/11/95	23,000	2,312	160	1,500	1.27 x 10 ⁵
09/13/95	12,000	532	10	230	1.84 x 10 ⁵
09/27/95	20,000	4,508	180	4,060	1.65 x 10 ³
10/19/95	12,000	3,854	120	2,960	1.21 x 10 ⁵
11/16/95	5,000	5,616	170	4,660	1.05 x 10 ⁵
12/19/95	20,000	1,920	60	1,520	1.19 x 10 ⁵
01/23/96	12,000	4,366	90	3,150	1.07 x 10 ⁵
02/13/96	12,000	11,232	110	6,510	0.90 x 10 ⁵

Lymphocyte subset counts in peripheral blood were determined by FACScan analysis as described previously (Ahmed-Ansari et al. [1989] Am. J. Primatol. 17:107-131).
 Virus levels were determined using the Chiron B-DNA assay as directed by the manufacturer. Plasma samples were stored

at - 80°C until use.

[°] ND, not done (no plasma sample was available for this date).

Table 2A. Alignment of HIV-1_{IC} (SEQ ID NO:2) and HIV-1_{NDK} env (SEQ ID NO:3) Protein Sequences. The consensus sequence corresponds to SEO ID NO:6. 1 MRvkEncqhlwrwgWKWGIMLLGMLMiCSAtEkLWVTVYYGVPvWKEtTTTLFCASDAKAY HIVJCENV 1 MRarE kerncqnlwKwGIMLLGMLMtCSAaEdLWVTVYYGVPiWKEaTTTLFCASDAKAY HIVNDKENV MR--En-----WKWGIMLLGMLM-CSA-E-LWVTVYYGVP-WKE-TTTLFCASDAKAY consensus HIVJCENV 62 eeEvHnvWaTHACVPTDPNPQEIvLaNVTEdFNMWKNeMVEQMHtDIISLWDeSLKPCVKL HIVNDKKNV 61 kkEaHNiWATHACVPTDPNPQEIeLeNVTEnFNMWKNnMVEQMHeDIISLWDqSLKPCVKL --E-HN-WATHACVPTDPNPQEI-L-NVTE-FNMWKN-MVEQMH-DIISLWD-SLKPCVKL consensus HIVJCENV 123 TPLCVTLNCTDlknEtktNSsdaNsnsgEimgnEeiKNCSFNVstgapgkvqkeYaLfyal 11 122 TPLCVTLNCTD eEkrKNCSFNVrdkreqvyalfYkL div HTVNDKENV E lrNS kgNgkveE TPLCVTLNCTD1knEt--NSs--N----Eimg-E--KNCSFNV------Y-Lf--consensus HIVJCENV 184 dlvsikNenNSTshmLtsCnTSvsTQACPKvSFEPIPIHyCAPAGFAILKCnDKKFNGTGP 174 pl dnnNrtNSTnyrLinCdTStiTQACPKiSFEPIPIHfCAPAGFAILKCrDKKFNGTGP HIVNDKENV -Iv---N--NST---L--C-TS--TQACPK-SFEPIPIH-CAPAGFAILKC-DKKFNGTGP consensus HIVJCENV 245 CnnvstvQCTHGIRPvvstQLLLNGsvAEEEvvlRSaNfsdNaKTIIVQLNhSveItCTRP Π HIVNDKENV 234 CSNVSTVQCTHGIRPVVSTQLLLNGS1AEEEiiiRSeNltnNvKTIIVQLNaSivInCTRP consensus C-NVSTVQCTHGIRPVVSTQLLLNGS-AEEE---RS-N---N-KTIIVQLN-S--I-CTRP **HIVJCENV** 306 nynetkkirIhrgygrsfvT vrKlGdrkQAHCtmnRtkWdnALkQiAsKLreqfNKTaI 1111 1 1 HIVNDKENV 295 ykytrqrtsIglrqslytiTgkkkKtGyigQAHCkisRaeWnkALqQvAtKLgnllNKTtI -----I----Tgk--K-G---QAHC---R--W--AL-Q-A-KL----NKT-I consensus ${\tt 365\ ifnrssGGDlEIemHsfnCGGelFYCNTtkLFNSTWNeTtesngkgeniTLPCRIrQfVNm}$ HIVJCENV 111111 | 11 HIVNDKENV 356 tFkpSSGGDpEItsHmlNCGGdfFYCNTsrLFNSTWNqTnstgfnngtvTLPCRIkQiVNl consensus -F--SSGGD-EI--H--NCGG--FYCNT--LFNSTWN-T-----TLPCRI-Q-VN-HIVJCENV 426 WQkVGKAMYAPPsdGqirCtSNITGLLLTRDGGhndNNtnnETfRPGrGDMRDNWRSELYK Π HIVNDKENV 417 WQrVGKAMYAPPieGlIkCsSNITGLLLTRDGG annsshetirPGgGDMRDNWRSELYK consensus WQ-VGKAMYAPP--G-I-C-SNITGLLLTRDGGhn-NN---ET-RPG-GDMRDNWRSELYK

Table 2A. (Continued)

HINJCENN	487	YKVİKIEPIGVAPTKAKRRVVQREKRAVGMVGAMFLGFLGAAGSTMGAASITLTVQARQLI
HIVNDKENV	476	YKVvKIEPIGVAPTKArRRVVEREKRAIG lGAvFLGFLGAAGSTMGAASVTLTVQARQLm
consensus		YKV-KIEP-GVAPTKA-RRVV-REKRA-Gm-GA-FLGFLGAAGSTMGAAS-TLTVQARQL-
HIVJCENV		SGIVqQQNNLLRAIEAQQHLLQLTVWGIKQLQARVLAVERYLKDQQLLGIWGCSGklICTT
HIVNDKENV	536	SGIVhQQNNLLRAIEAQQHLLQLTVWGIKQLQARVLAVERYLrDQQLLGIWGCSGrhICTT
consensus		SGIV-QQNNLLRAIEAQQHLLQLTVWGIKQLQARVLAVERYL-DQQLLGIWGCSGICTT
HIVJCENV	609	tVPWNaSWSNkSLDqIWnNMTWmEWdREIaNYTnLIhhLIEESQnQQEKNEqELLELDKWA
HIVNDKENV	597	nvPwnsSwSnrsldeiwqnmTwmEweREIdNYTgLIysLIEESQiQQEKNEkELLELDKWA
consensus		-VPWN-SWSN-SLD-IW-NMTW-EW-REI-NYT-LILIEESQ-QQEKNE-ELLELDKWA
HIVJCENV	670	SLWsWFdlsnWLWYIKiFIMIVaGLvGLRIVFAVLSiVNRVRQGYSPLSFQThfPaPRGPD
HIVNDKENV	658	SLWnWFsitkWLWYIKlFIMIVgGLiGLRIVFAVLsvVNRVRQGYSPLSFQTllPvPRGPD
consensus		SLW-WF-IWLWYIK-FIMIV-GL-GLRIVFAVLS-VNRVRQGYSPLSFQTP-PRGPD
HIVJCENV	731	RPdgIEgEGGERdRDRSvRLVdGflALlWeDLRNLCLFSYHRLRDllLIvtRIVELLGRRG
HIVNDKENV	719	RPeeIEeEGGERGRDRSiRLVnGlfALfWdDLRNLCLFSYHRLRDsiLIaaRIVELLGRRG
consensus		RPIE-EGGER-RDRS-RLV-GAL-W-DLRNLCLFSYHRLRDLIRIVELLGRRG
HIVJCENV	792	WEALKYLWSLLQYWSQELkNSAVnLfnTtAIvVAEgTDRiIEVVQRlCRAILhiPRRIRQG
HIVNDKENV	780	WEALKYLWnLLQYWSQELrNSAsslldTiAIaVAErTDRvIEVVQRaCRAILnvPRRIRQG
consensus		WEALKYLW-LLQYWSQEL-NSALT-AI-VAE-TDR-IEVVQR-CRAILPRRIRQG
HIVJCENV	853	LERILL
HIVNDKENV	841	LERILL
consensus		LER-LL

Alignment score = 541.00

 $\label{eq:table 2B.} \textbf{Alignment of HIV-1}_{\text{IC}} \ \text{and HIV-1}_{\text{LAI}} \ \underline{\text{env}} \ \text{Protein Sequences (SEQ ID NO:2 and SEQ ID NO:4, respectively)}. \ \ \text{The consensus sequence corresponds to SEQ ID NO:7} \ .$

HIVJCENV	1	${\tt MRVKEncQHLWRWGWKWGiMLLGmLMICSATEKLWVTVYYGVPVWKBtTTTLFCASDAKA} {\tt MRVKEncQHLWRWGWKWGimlGmLMICSATEKLWVTVYYGVPVWKBtTTTLFCASDAKA} {\tt MRVKEncQHLWRWGWKWGimlGmLMICSATEKLWVTVYYGVPVWKBtTTTLFCASDAKA} {\tt MRVKEncQHLWRWGWKWGimlGmLMICSATEKLWVTVYYGVPVWKBtTTTLFCASDAKA} {\tt MRVKEncQHLWRWGWKWGimlGmLMICSATEKLWVTVYYGVPVWKBtTTTLFCASDAKA {\tt MRVKEncQHLWRWGWKWGimlGmLMICSATEKLWVTVYYGVPVWKBTTTTLFCASDAKA {\tt MRVKEncQHLWRWGWKWGimlGmLMICSATEKLWVTVYYGVPVWKBTTTTLFCASDAKA {\tt MRVKEncQHLWRWGWKWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmLWGimlGmlW$
HIVlaienvt	1	MRVKEkyQHLWRWGWKWGtMLLGiLMICSATEKLWVTVYYGVPVWKEaTTTLFCASDAKA
consensus		MRVKEQHLWRWGWKWG-MLLG-LMICSATEKLWVTVYYGVPVWKE-TTTLFCASDAKAY
HIVJCENV	62	eeEVHNVWATHACVPTDPNPQEiVLaNVTEdFNMWKNeMVEQMHtDIISLWDeSLKPCVKI
HIVlaienvt	62	dtevhnvwathacvptdpnpqsvvlvnvtenfnmwkndmveqmhediislwdqslkpcvki
consensus		EVHNVWATHACVPTDPNPQE-VL-NVTE-FNMWKN-MVEQMH-DIISLWD-SLKPCVKI
HIVJCENV	123	TPLCVtLnCTDLkneTkTNSSdaNSnSGEiM gneEIKNCSFNvSTgapGKVQKEYalFYa
HIVlaienvt	123	TPLCVsLkCTDLgNaTnTNSSntNSsSGEmMmekgEIKNCSFNiSTsirGKVQKEYafFYk
consensus		TPLCV-L-CTDL-N-T-TNSSNS-SGE-MmEIKNCSFN-STGKVQKEYFY-
HIVJCENV	183	LDIvsIkneNnsTShmLTSCNTSVsTQACPKVSFEPIPIHYCAPAGFAILKCNdKkFNGTG
HIVlaienvt	184	LDI IpidNdtTSytLTSCNTSViTQACPKVSFEPIPIHYCAPAGFAILKCNnKtFNGTG
consensus		LDIvsINTSLTSCNTSV-TQACPKVSFEPIPIHYCAPAGFAILKCN-K-FNGTG
HIVJCENV	244	PCnnvstvQcthgirpvvstQLLLngsvaEEEvvlrsanfsDnaktiivQLnhsvEitcTr
HIVlaienvt	243	PCtnVsTVQCTHGIRPVVsTQLLLNGSlaEEEVViRSANFtDNAKTIIVQLNqSVBInCTF
consensus		PC-NVSTVQCTHGIRPVVSTQLLLNGS-AEEEVV-RSANF-DNAKTIIVQLN-SVEI-CTR
HIVJCENV		PNyNetKkIRIhRGyGRsFVTvrKlGdrkQAHCtmnRtKWdnaLKQIASKLREQFnNktaI
HIVlaienvt		PNnNtrKsIRIqRGpGRaFVTigKiGnmrQAHCnisRaKWnatLKQIASKLREQFgNnktI
consensus		PN-NK-IRI-RG-GR-FVTK-GQAHCR-KWLKQIASKLREQFgNI
HIVJCENV	365	IFnrssggDleIemHsfncggelfycnttkLfnstw net te sn gkg enItLpcrI
IIVlaienvt	365	IFkqSSGGDpEIvtHSFNCGGEfFYCNsTqLFNSTWfNsTwsTEgSNnteGsdtITLPCRI
consensus		IFSSGGD-EIHSFNCGGE-FYCN-T-LFNSTWfN-TwsTEgSNnGsITLPCRI
HIVJCENV	420	rQFvNMWQkVGKAMYAPPsdGQIRCtSNITGLLLTRDGGhNdNNtnnEtFRPGrGDMRDNW
IIVlaienvt	426	kQFiNMWQeVGKAMYAPPisGQIRCsSNITGLLLTRDGG N NNngsEiFRPGGGDMRDNW
		OF MESO VOVINCED COTTO CUTTOUT CONTROLL STORY OF FREE CHICAGO

Table 2B. (Continued)

HIVJCENV	481 RSELYKYKVİKIEPLGVAPTKAKRRVVQREKRAVGMVGAMFLGFLGAAGSTMGAASİTLTV
consensus	RSELYKYKV-KIEPLGVAPTKAKRRVVQREKRAVGm-GA-FLGFLGAAGSTMGA-S-TLTV
HIVJCENV	542 QARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARvLAVERYLKDQQLLGIWGCSG
HIVlaienvt	545 QARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARiLAVERYLKDQQLLGIWGCSG
consensus	QARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQAR-LAVERYLKDQQLLGIWGCSG
HIVJCENV	603 KLICTTtVPWNASWSNKSLdQIWNNMTWmEWDREIaNYTnLIHhLIEESQNQQEKNEQELL
:	
HIVlaienvt	606 KLICTTaVPWNASWSNKSLeQIWNNMTWmEWDREInNYTsLIHsLIEESQNQQEKNEQELL
consensus	KLICTT-VPWNASWSNKSL-QIWNNMTW-EWDREI-NYT-LIH-LIEESQNQQEKNEQELL
HIVJCENV	664 ELDKWASLWsWFdIsNWLWYIKIFIMIVAGLVGLRIVFAVLSIVNRVRQGYSPLSFQTHfP
HIVlaienvt	
consensus	ELDKWASLW-WF-I-NWLWYIKIFIMIV-GLVGLRIVFAVLSIVNRVRQGYSPLSFQTH-P
HIVJCENV	725 aPRGPDRPdGIEgEGGERDRDRSvRLVdGfLALlWeDLRnLCLFSYHRLRDLLLIVTRIVE
HIVlaienvt	728 tPRGPDRPeGIEeEGGERDRDRSiRLVnGsLALiWdDLRsLCLFSYHRLRDLLLIVTRIVE
consensus	-PRGPDRP-GIE-EGGERDRDRS-RLV-G-LAL-W-DLR-LCLFSYHRLRDLLLIVTRIVE
HIVJCENV	786 LLGRRGWEALKYlWsLLQYWSQELKNSAVnLfNtTAIvVAEGTDRiIEVVQrlCRAIlHIP
III VUCLIVV	
HIVlaienvt	789 LLGRRGWEALKYWWnLLQYWSQELKNSAVsLlNaTAIaVAEGTDRvIEVVQgaCRAIrHIP
consensus	LLGRRGWEALKY-W-LLQYWSQELKNSAV-L-N-TAI-VAEGTDR-IEVVQCRAI-HIP
11 7 11 7/1030 *	947 PRIDOCIERIII
HIVJCENV	847 RRIRQGLERILL
HIVlaienvt	850 RRIRQGLERILL
consensus	RRIRQGLER-LL

Alignment score = 657.00

Table 2C. Alignment of HIV-1_{IC} and HIV-1_{SF2} env Protein Sequences (SEQ ID NO:2 and SEQ ID NO:5, respectively). The consensus sequence corresponds to SEQ ID NO:8.

HIVJCENV	1 MrVKencqhlwrwgWkWGimLLGMLMICSATEKLWVTVYYGVPVWKEtTTTLFCASDAkAY
HIVsf2envt	1 MkVK gtrrnyqhlwrwGtlLLGMLMICSATEKLWVTVYYGVPVWKEaTTTLFCASDArAY
consensus	M-VKeW-WGLLGMLMICSATEKLWVTVYYGVPVWKE-TTTLFCASDA-AY
HIVJCENV	62 eeEVHNVWATHACVPTDPNPQEiVLaNVTEdFNMWKNeMVEQMhtDIISLWDeSLKPCVKL
HIVsf2envt	61 dtevenvwathacvptdpnpqevvlgnvtenfnmwknnmveqmqedIIsLwdqSLkPCVkL
consensus	EVHNVWATHACVPTDPNPQE-VL-NVTE-FNMWKN-MVEQMDIISLWD-SLKPCVKL
HIVJCENV	123 TPLCVTLNCTDLkneTkTNSSdaNsnsgeimgneeikncsfnvstgapgKvqkeyalfyal
HIVsf2envt	122 TPLCVTLNCTDLgkaTnTNSS NwkeeikgeikncsfnittsirdkiqKenalfrnldvv
consensus	TPLCVTLNCTDLT-TNSSdaNKKK
HIVJCENV	184 dlvsiknenNsTshmLtsCNtSVsTQACPKVSFEPIPIHYCaPAGFAILKCNdKkFNGtGP
HIVsf2envt	181 pIdnastttNyTnyrLihCNrSViTQACPKVSFEPIPIHYCtPAGFAILKCNnKtFNGkGP
consensus	-IN-TLCN-SV-TQACPKVSFEPIPIHYC-PAGFAILKCN-K-FNG-GP
HIVJCENV	245 CnnvstvQCTHGIRPvvstQLLLnGsvAEEEvvlRsanfsdnAkTIIvQLnhsVeltCTRP
HIVsf2envt	242 CtnvstvQCTHGIRPivstQLLLNGSlAFEEvvirsdnFtnNAKTIIvQLNesVaInCTRP
consensus	C-NVSTVQCTHGIRP-VSTQLLLNGS-ABEEVV-RS-NFNAKTIIVQLN-SV-I-CTRP
HIVJCENV	306 NyNetKkIrIhrGygrsfvTvRklGDrkqAHCtmnRtkWdNaLkQIasKLREQFnNktaIi
HIVsf2envt	303 NnNtrKslylgpG rafhtTgRiiGDirkAHCnisRaqWnNtLeQlvkKLREQFgNnktlv
consensus	N-NK-I-IGyT-RGDAHCRW-N-L-QIKLREQFgNI-
HIVJCENV	366 FNrssGGDlEIeMHSFNCgGElFYCNTTkLFNsTW netTEsngkgenItLPCRIrQfvNM
HIVsf2envt	
consensus	FN-SSGGD-EI-MHSFNC-GE-FYCNTT-LFN-TWrTEI-LPCRI-QNM
	426 WQKVGKAMYAPPadGQIrCtSNITGLLLTRDGGhNdnNtnnEtFRPGrGDMRDNWRSELYK
consensus	WO-VGKAMYAPPGOI-C-SNITGLLLTRDGG-NNtE-FRPG-GDMRDNWRSELYK

Table 2C. (Continued)

HIVJCENV	487	YKVIKIEPLGvAPTKAKRRVVQREKRAVGmVGAMFLGFLGAAGSTMGAASLTLTVQARQLL
HIVsf2envt	484	
consensus		YKVIKIEPLG-APTKAKRRVVQREKRAVG-VGAMFLGFLGAAGSTMGA-SLTLTVQARQLL
HIVJCENV	548	SGIVQQQNNLLRAIRAQQHLLQLTVWGIKQLQARVLAVERYLkDQQLLGIWGCSGKLICTT
HIVsf2envt	545	SGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARVLAVERYL±DQQLLGIWGCSGKLICTT
consensus		SGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARVLAVERYL-DQQLLGIWGCSGKLICTT
HIVJCENV	609	tVPWNASWSNKSLdqIWnNMTWmeWdREIANYTNlIhhLiEESQNQQEKNEQELLELDKWA
HIVsf2envt	606	aVPWNASWSNKSLediwdnmTwmqWerEidnYTntiytllEESQNQQEKNEQELLELDKWA
consensus		-VPWNASWSNKSLIW-NMTWW-REI-NYTN-IL-BESQNQQEKNEQELLELDKWA
HIVJCENV	670	SLWsWFdIsnwLwyIKIFIMIVaGLVGLRIVFAVLSIVNRVRQGYSPLSFQThfPaPRGPD
HIVsf2envt	667	SLWnWFsitnWLWYIKIFIMIVgGLVGLRIVFAVLSIVNRVRQGYSPLSFQTrlpvPRGPD
consensus		SLW-WF-I-NWLWYIKIFIMIV-GLVGLRIVFAVLSIVNRVRQGYSPLSFQTP-PRGPD
HIVJCENV	731	RPDGIEGEGGERDRDRSVRLVDGFLALlWEDLRnLCLFSYhRLRDLLLIvtRiVElLGrRG
HIVsf2envt	728	RPDGIEeEGGERDRDRSVRLVDGFLALiWEDLRsLCLFSYrRLRDLLLIaaRtVEiLGhRG
consensus		RPDGIE-EGGERDRDRSVRLVDGFLAL-WEDLR-LCLFSY-RLRDLLLIR-VE-LG-RG
HIVJCENV	792	WEALKYlWSLLQYWSQELKNSAVnlfntTAIvVaEGTDRiIEVvQRlcRAILHIPRRIRQG
HIVsf2envt	789	WEALKYwWSLLQYWiQELKNSAVswlNaTAIaVtEGTDRvIEVaQRayRAILHIhRRIRQG
consensus		WEALKY-WSLLQYW-QELKNSAVN-TAI-V-EGTDR-IEV-QRRAILHI-RRIRQG
HIVJCENV	853	LERILL
HIVsf2envt	850	LERILL
consensus		LER-LL
Alignment s	core	- 614 00

Table 3. HIV-1 Antibody Response of C534 After Transfusion From C455

Days Post Infection	Titer
0	0
7	0
. 14	400
21	1600
28	1600

Table 4. Comparative Analyses of Chimpanzee HIV-1 Isolates With Inoculating Viruses

			HIV-1 _{JC}	16 vs:		Mutations
		LAI	SF2	NDĶ	NC7	unique to JC/NC ²
Pol	% homology ¹ insertions deletions point mutations	96.6 12 0 30	94.8 0 0 51	94.0 0 1 56	98.8 0 0 11	0 0 23
Gag	% homology ¹ insertions deletions point mutations	94.2 12 0 25	92.2 5 3 33	88.8 1 81 42	97.4 2 0 9	- 0 2 20
Vif	% homology ¹ insertions deletions point mutations	92.2 0 0 14	86.0 0 0 26	86.5 0 0 25	96.9 0 0 5	- 0 0 10
Vpr	% homology ¹ insertions deletions point mutations	88.7 0 1 7	96.9 0 0 2	89.7 0 1 6	96.6 0 0 2	- 0 0
Tat	% homology ¹ insertions deletions point mutations	88.4 0 15 10	82.5 1 0 18	74.4 0 15 22	97.0 0 0 3	0 0 4
Nef	% homology ¹ insertions deletions point mutations	87.4 0 0 25	89.4 4 0 18	78.7 1 0 40	99.0 0 .0 1	- 0 0 8
Rev	% homology ¹ insertions deletions point mutations	84.5 1 1 18	82.9 1 0 19	75.2 1 0 28	96.6 0 0 4	- 0 0 11
Env	% homology ¹ insertions deletions point mutations	81.9 9 7 138	76.1 9 13 173	71.4 11 24 203	97.6 0 1 17	- 0 2 78
Vpu	% homology ¹ insertions deletions point mutations	79.3 0 1 13	63.9 0 1 24	67.1 0 1 24	98.8 0 0 0	- 0 0 10
LTR	% homology ¹ insertions deletions point mutations	92.1 1 1 48	93.9 3 3 29	89.1 1 1 52	98.7 0 0 10	0 1 14

Amino acid homology
Amino acids or nucleotides not observed in parental viruses
Nucleotide identity

MBD -> matrix protein (p17)	GCRQVLEQLQPSLQT 70I.G 70I.G 70
GSEELR GSEELR Capsid I IVONLO TINEEA	
9a9 9a9 5 gag GSEELR 9a9 9a9 5 gag IVONLO 9a9 9a9 9a9 9a9	
9a9 9 gag GSEELR 9a9 9a9 9a9 9a9 9a9 9a9 9a9 9a9 9a9 9a9	0/
6 gag GSEELR gag	K.IIG1., 70
GSEELR capsid IVONLO IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII .	 FEL GANOSADBABTO
9449 9449 9449 9449 9449 9449 9449 944	
	133
pagI. capsid pag IVQNLQ pag roll roll roll roll pag roll TGN 135	
9 9 9 1 N E E A 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	
Jag	
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н н н н н н н н н н н н н н н н н н н	6/2
H	272
	270
C dad V	DCKTILKALGPGATL 343
:	
	343 A
HIV-1SF2 gagT	040A
:	
HIV-1JC16 gag EEMMTACQGVGGPGHKARVLABAMSQVT-NSAAIMMQRGNFKNQRKMVKCFNCGKEGHVARNCRAPRKKG	
	Δ13 · · · · · · · · · · · · · · · · · · ·
HIV-ILAI gag	H . K
HIV-ISFZ gagT.T.R.T.R.T.N.T.NDK gagT.R.T.R.T.R.T.R.T.R.T.R.T.R.T.R.T.	T.K410

470	472	482	472	420		501	503	513	503	420
finger) p7 <- p1	:	:								
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SG	:	•	ъ.							
LR	:	[24	<u>Γτ.</u>							
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!	:	FLO	1							
d.	•	:	•							
A T	:	:	:							
7 13	:	:	:							
p6 SR	:	:	•							
- 310	:	:	:							
{ zinc finger } p7 <- p1 -> p6 CWKCGKEGHQMKDCIESQANFLGKIWPSHKGRPGNFLQSR		T.R Y	:							
RP	:	:	:							
 H K G	•		×	•	-	œ	:	:	:	
-p1 PS1	:	•	•		-> 9d	PS	:	:	:	
	:	•	:			ONE	:	:	:	
	:	:	:			LFC				
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S		:	:	:		T P	•			
zinc] HIV-1JC16 qaq CWKCGF	HIV-1NC7 dad	ם מ	a d	HIV-1NDK gagR		HIV-1JC16 qaq TPSQKQEPIDKEVYPLTSLRSLFGNDPSSQ	a d	HIV-1LAI gag	ad	ag
16 0	7) 6 H	ָ מו	1 K	1	91	7	ים א	0	HIV-1NDK gag
1.30	INC	1I.A	1SF	1ND		1,70	INC	IIA	1SF	1 ND
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Table 5 (continued)

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MTD		
HIV-1JC16 nef MGGKF	HIV-1JC16 nef MGGKWSKSSIVGWPTIRERMKRAGPAADGVGAASRDLEKHGAITSSNTAATNADCA	26
HIV-1NC7 nef		99
HIV-1LAI nef		26
HIV-1SF2 nef	R.MG.SAR.BPRAEVV	9
HIV-1NDK nef		26
	LCK binding domain	
HIV-1JC16 nef WLEAC	HIV-1JC16 nef WLEAQEE-BEVGFPVRPQVPLRPMTYKAGIDLSHFLKEKGGLEGLVWSQRRQDILDLWIY 115	115
HIV-1NC7 nef	1918	115
HIV-1LAI nef	TAVAV	115
HIV-1SF2 nef		119
HIV-1NDK nef	S	116
HIV-1JC16 nef HTQG1	HIV-1JC16 nef HTQGYFPDWQNYTPGPGIRYPLTFGWCFKLVPVEPDKVBEANBGBDNILLHPMCLHGMBD 175	175
HIV-1NC7 nef		175
HIV-1LAI nef		175
HIV-1SF2 nef		179
HIV-1NDK nef N1	$N\dots I\dots \dots \dots \dots \dots Q\dots \dots Q\dots \dots D \dots Q \dots \dots \dots \dots \dots \dots \dots $	176
HIV-1JC16 nef AEKEV	HIV-1JC16 nef AEKEVLVWRFDSKLAPHHVARBLHPBYYKDC	207
HIV-1NC7 nef		207
HIV-1LAI nef P.R.	D. R E	207
HIV-1SF2 nef		211
HTV-1NDK nef P RO	HTV-1NDK nof D RO M . N R I E K	208

Table 5 (continued)		
	signal peptide <- ->gp120 (SU)	
6 env MRVKE-	NCQHLWRWGWKWGIMLLGMLMICSATEKLWVTVYYGVPVWKETTTTLFCASD 57	
HIV-1NC7 envY.	57	
HIV-1LAI envKY.	I	
HIV-1SF2 env . K GTRR . Y		
HIV-1NDK env AR.KBR	AR.KBRNTA.DIA.56	
HIV-1JC16 env AKAYBEBVHN	AKAYBEBVHNVWATHACVPTDPNPQEIVLANVTEDFNMWKNEMVEOMHTDIISLWDBSLK 117	_
HIV-1NC7 env		. ~
HIV-1LAI envDT		. ~
HIV-1SF2 env . R DT		
HIV-1NDK envKK.A	B 0	
env	PCVKLTPLCVTLNCTDLKNETKTNSSDANSNSGEIM-GNEEIKNCSFNVSTGAPGKVQKE 176	
env	176	ı^
HIV-1LAI env	S.KG.A.NNTSM.MEKGISIR 177	_
HIV-1SF2 env		_
HIV-1NDK env		_
	V2	
HIV-1JC16 env YALFYALDIVS	YALFYALDIVSIKNENNSTSHMLTSCNTSVSTQACPKVSFEPIPIHYCAPAGFAILK 233	_
env	233	_
HIV-1LAI envFKII	P.D.DTT YT I 1 232	٠,
HIV-1SF2 env N RN V . I		_
HIV-1NDK envK	KP.D.N.RTNTNYR.IN.DTIIF 222	61
HIV-1JC16 env CNDKKFNGTG1	CNDKKFNGTGPCNNVSTVQCTHGIRPVVSTQLLLNGSVAEEEVVLRSANFSDNAKTIIVQ 293	_
HIV-1NC7 env		_
env	T	61
HIV-1SF2 envN.TK.	N . T K T	_
HIV-1NDK env . R	RB.LTN.V 282	61
env	SYGRSFVTV RKLGDRKQAHCTMNRTKWDNALK	•
env		Φ.
env	-KINMRNIS.ANAT	~
env	N.TR.S.Y .PA.H.TGIIIRKNIS.AQ.N.T.E 345	ıc
HIV-1NDK env . A. IV. N.		_

Table 5 (continued)		
HIV-1JC16 env	OIASKLREOFNNKTAIIFNRSSGGDLEIEMHSFNCGGELFYCNTKLFNSTWNR 403	~
HIV-1NC7 env		ואו
HIV-1LAI env		
HIV-1SF2 env	VKG.NKT.VQPVRFQNWRLN 400	0
HIV-1NDK env	.V.TGNLLT.T.KPPTS.MLDFSRQ 393	۳
	CD4 binding	
HIV-1JC16 env	TTESNGKGENITLPCRIROPVNMWOKVGKAMYAPPSDGOIRCTSNITGLLLTRD	~
HIV-1NC7 env	OSAB.	. ~
HIV-1LAI env		7
HIV-1SF2 env	H. GTKGNDT.I K.II E IG S.S T.VT 460	0
HIV-1NDK env	.NSTGFNNGTVK.ILRIB.L.K.SA 451	-
	V5 gp120 (SU) <- -> gp41 (TM)	
HIV-1JC16 env	_	e
HIV-1NC7 env	. KTDKG	7
HIV-1LAI env	GS.IG	ហ
HIV-1SF2 env	DT.VGII	6
HIV-1NDK env	$. ssh I\ldots g\ldots$	0
HIV-1JC16 env	GFLGAAGSTMGAASLTLTVOAROLLSGIVOOONNLLRAIBAOOHLLOLTVWGIKOLOARV 583	~
HIV-1NC7 env		. ~
		ស
HIV-1SF2 env	979VV	6
HIV-1NDK env		0
HIV-1JC16 env	HIV-1JC16 env LAVBRYLKDQQLLGIWGCSGKLICTTTVPWNASWSNKSLDQIWNNMTWMEWDREIANYTN 643	m
HIV-1NC7 env	645	7
HIV-1LAI env	849 S N	ហ
HIV-1SF2 env	R	6
HIV-1NDK env	$\dots \dots $	0
HIV-1JC16 env	LIHHLIBESQNQQEKNBQELLELDKWASLWSWFDISNWLWYIKIFIMIVAGLVGLRIVFA 703	ú.
HIV-1NC7 env		Ņ
HIV-1LAI env	s	ñ
HIV-1SF2 env	T.YT.L	ō,
HIV-1NDK env	YSIKKNS.TKLGI 690	0

Table 5 (continued)

763	765	759	750	823	822	825	819	810	960	859	862	3 2 8	244	5
REDEBERS VRIVDG FLALLWEDL	IN.SI.D	· · · · I · · · · · · · · · · · · · · ·	GIN.LFF.D	LLQYWSQELKNSAVNLFNTTAI	a	I. S. I. S	A. TAS I	I						
HIV-1JC16 env VLSIVNRVRQGYSPLSFQTHFPAPRGPDRFDGIRGEGGERDRDRSVRLVDGFLALLWEDL 763 HIV-1NC7 env		RL.V	HIV-INDK envVLFL.VEEEGIN.LFF.D 750	HIV-1JC16 env RNLCLFSYHRLRDLLLIVTRIVELLGRRGWEALKYLWSLLQYWSQELKNSAVNLFNTTAI 823		HIV-1LAI env SS.IL.B.I.	. S R AA.T I H		HIV-1JC16 env VVABGTDRIIEVVQRLCRAILHIPRRIRQGLERILL	HIV-1NC7 envF	HIV-1LAI env AVGARR	HIV-1SF2 env A.TVAAYHL	HIV-1NDK env ARVANVL	
env		env	env	env	env	env	env		env	env	env	env	env	
HIV-1JC16 env HIV-1NC7 env	HIV-1LAI env	HIV-1SF2 env	HIV-INDK	HIV-1JC16	HIV-1NC7 env	HIV-1LAI	HIV-1SF2 env	HIV-1NDK env	HIV-1JC16	HIV-1NC7	HIV-1LAI	HIV-1SF2	HIV-1NDK	

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: EMORY UNIVERSITY
 - (ii) TITLE OF INVENTION: Human Immunodeficiency Viruses
 Causing AIDS in a Nonhuman Primate
 - (iii) NUMBER OF SEQUENCES: 23
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Greenlee, Winner and Sullivan, P.C.
 - (B) STREET: 5370 Manhattan Circle, Suite 201
 - (C) CITY: Boulder
 - (D) STATE: Colorado
 - (E) COUNTRY: US
 - (F) ZIP: 80303
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: WO
 - (B) FILING DATE: 23-JUN-1998
 - (C) CLASSIFICATION:
 - (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 60/050,548
 - (B) FILING DATE: 23-JUN-1997
 - (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 60/057,606
 - (B) FILING DATE: 04-SEP-1997
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Ferber M., Donna
 - (B) REGISTRATION NUMBER: 33,878
 - (C) REFERENCE/DOCKET NUMBER: 66-97 WO
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (303) 499-8080
 - (B) TELEFAX: (303) 499-8089
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2577 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: not relevant

• WQ 98/59074 - PCT/US98/12990

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: 1..2577

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

	GTG Val	_						48
	ATC Ile							96
	TGG Trp 35							144
	ACT Thr							192
	AAT Asn							240
	GAA Glu							288
	GAA Glu							336
	AGC Ser 115							384
	TGC Cys							432
	AGT Ser							480
	TTC Phe							528

- W(98/5	9074			-						PCT/U	S98/12990
				Tyr			Val				AAT Asn	576
			Ser								ACA Thr	624
			CCA Pro									672
			GGT Gly									720
			CCA Pro									768
			GTA Val 260									816
			GTA Val									864
			GTA Val									912
			AAT Asn									960
			GTT Val									1008
			AAT Asn 340									1056
			AGA Arg									1104
TCC Ser			GGG Gly									1152
GGG Gly 385										Thr		1200

. w	98/5	9074		-					PCT/US98	/129 9 0
-			ACA Thr						CCA Pro	1248
			AGA Arg 420						GCA Ala	1296
			CCT Pro					 	 	1344
			CTA							1392
			ACC Thr							1440
			TTA Leu							1488
			ACC Thr 500							1536
			ATG Met							1584
			ATG Met							1632
			TCT Ser							1680
			CAA Gln							1728
			GCA Ala 580							1776
			GGG Gly							1824
			TGG Trp							1872

- W(98/5	9074						-		PCT/US98/	12990
					TGG Trp 630						1920
					CAC His						1968
					TTA Leu						2016
				_	TCA Ser						2064
					TTA Leu						2112
					GTT Val 710						2160
					CCG Pro						2208
					AGA Arg						2256
			_		CTC Leu						2304
					AGA Arg						2352
					AGG Arg 790						2400
					AGT Ser						2448
					ATA Ile						2496
					CTT Leu						2544

ATT AGA CAG GGC TTG GAA AGA TTT TTG CTA TAA
Ile Arg Gln Gly Leu Glu Arg Phe Leu Leu *
850

2577

- (2) INFORMATION FOR SEQ ID NO:2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 859 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
- Met Arg Val Lys Glu Asn Cys Gln His Leu Trp Arg Trp Gly Trp Lys
- Trp Gly Ile Met Leu Leu Gly Met Leu Met Ile Cys Ser Ala Thr Glu 20 25 30
- Lys Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Val Trp Lys Glu Thr
 35 40 45
- Thr Thr Thr Leu Phe Cys Ala Ser Asp Ala Lys Ala Tyr Glu Glu Glu 50 55 60
- Val His Asn Val Trp Ala Thr His Ala Cys Val Pro Thr Asp Pro Asn
 65 70 75 80
- Pro Gln Glu Ile Val Leu Ala Asn Val Thr Glu Asp Phe Asn Met Trp 85 90 95
- Lys Asn Glu Met Val Glu Gln Met His Thr Asp Ile Ile Ser Leu Trp
 100 105 110
- Asp Glu Ser Leu Lys Pro Cys Val Lys Leu Thr Pro Leu Cys Val Thr 115 120 125
- Leu Asn Cys Thr Asp Leu Lys Asn Glu Thr Lys Thr Asn Ser Ser Asp 130 135 140
- Ala Asn Ser Asn Ser Gly Glu Ile Met Gly Asn Glu Glu Ile Lys Asn 145 150 155 160
- Cys Ser Phe Asn Val Ser Thr Gly Ala Pro Gly Lys Val Gln Lys Glu 165 170 175
- Tyr Ser Leu Phe Tyr Ala Leu Asp Ile Val Ser Ile Lys Asn Glu Asn 180 185 190
- Asn Ser Thr Ser His Met Leu Thr Ser Cys Asn Thr Ser Val Ser Thr 195 200 205
- Gln Ala Cys Pro Lys Val Ser Phe Glu Pro Ile Pro Ile His Tyr Cys 210 215 220

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Ala	Pro	Ala	Gly	Phe	Ala	Ile	Leu	Lys	Cys	Asn	Asp	Lys	Lys	Phe	Asn
225.					230					235					240

- Gly Thr Gly Pro Cys Asn Asn Val Ser Thr Val Gln Cys Thr His Gly 245 250 255
- Ile Arg Pro Val Val Ser Thr Gln Leu Leu Leu Asn Gly Ser Val Ala
 260 265 270
- Glu Glu Glu Val Val Leu Arg Ser Ala Asn Phe Ser Asp Asn Ala Lys 275 280 285
- Thr Ile Ile Val Gln Leu Asn His Ser Val Glu Ile Thr Cys Thr Arg
- Pro Asn Tyr Asn Glu Thr Lys Lys Ile Arg Ile His Arg Gly Tyr Gly 305 310 315 320
- Arg Ser Phe Val Thr Val Arg Lys Leu Gly Asp Arg Lys Gln Ala His 325 330 335
- Cys Thr Met Asn Arg Thr Lys Trp Asp Asn Ala Leu Lys Gln Ile Ala 340 345 350
- Ser Lys Leu Arg Glu Gln Phe Asn Lys Thr Ala Ile Ile Phe Asn Arg 355 360 365
- Ser Ser Gly Gly Asp Leu Glu Ile Glu Met His Ser Phe Asn Cys Gly 370 375 380
- Gly Glu Leu Phe Tyr Cys Asn Thr Thr Lys Leu Phe Asn Ser Thr Trp 385 390 395 400
- Asn Glu Thr Thr Glu Ser Asn Gly Lys Gly Glu Asn Ile Thr Leu Pro . 405 410 415
- Cys Arg Ile Arg Gln Phe Val Asn Met Trp Gln Lys Val Gly Lys Ala
 420 425 430
- Met Tyr Ala Pro Pro Ser Asp Gly Gln Ile Arg Cys Thr Ser Asn Ile 435 440 445
- Thr Gly Leu Leu Thr Arg Asp Gly Gly His Asn Asp Asn Asn Thr 450 455 460
- Asn Asn Glu Thr Phe Arg Pro Gly Arg Gly Asp Met Arg Asp Asn Trp 465 470 475 480
- Arg Ser Glu Leu Tyr Lys Tyr Lys Val Ile Lys Ile Glu Pro Leu Gly
 485 490 495
- Val Ala Pro Thr Lys Ala Lys Arg Arg Val Val Gln Arg Glu Lys Arg 500 505 510
- Ala Val Gly Met Val Gly Ala Met Phe Leu Gly Phe Leu Gly Ala Ala 515 520 525

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- Gly Ser Thr Met Gly Ala Ala Ser Leu Thr Leu Thr Val Gln Ala Arg 530 540
- Gln Leu Leu Ser Gly Ile Val Gln Gln Gln Asn Asn Leu Leu Arg Ala 545 550 555 560
- Ile Glu Ala Gln Gln His Leu Leu Gln Leu Thr Val Trp Gly Ile Lys
 565 570 575
- Gln Leu Gln Ala Arg Val Leu Ala Val Glu Arg Tyr Leu Lys Asp Gln
 580 585 590
- Gln Leu Leu Gly Ile Trp Gly Cys Ser Gly Lys Leu Ile Cys Thr Thr 595 600 605
- Thr Val Pro Trp Asn Ala Ser Trp Ser Asn Lys Ser Leu Asp Gln Ile 610 615 620
- Trp Asn Asn Met Thr Trp Leu Glu Trp Asp Arg Glu Ile Ala Asn Tyr 625 630 635 640
- Thr Asn Leu Ile His His Leu Ile Glu Glu Ser Gln Asn Gln Glu 645 650 655
- Lys Asn Glu Gln Glu Leu Leu Glu Leu Asp Lys Trp Ala Ser Leu Trp 660 665 670
- Ser Trp Phe Asp Ile Ser Asn Trp Leu Trp Tyr Ile Lys Ile Phe Ile 675 680 685
- Met Ile Val Ala Gly Leu Val Gly Leu Arg Ile Val Phe Ala Val Leu 690 695 700
- Ser Ile Val Asn Arg Val Arg Gln Gly Tyr Ser Pro Leu Ser Phe Gln 705 710 715 720
- Thr His Phe Pro Ala Pro Arg Gly Pro Asp Arg Pro Asp Gly Ile Glu
 725 730 735
- Gly Glu Gly Glu Arg Asp Arg Asp Arg Ser Val Arg Leu Val Asp 740 745 750
- Gly Phe Leu Ala Leu Leu Trp Glu Asp Leu Arg Asn Leu Cys Leu Phe 755 760 765
- Ser Tyr His Arg Leu Arg Asp Leu Leu Leu Ile Val Thr Arg Ile Val
 770 780
- Glu Leu Leu Gly Arg Arg Gly Trp Glu Ala Leu Lys Tyr Leu Trp Ser 785 790 795 800
- Leu Leu Gln Tyr Trp Ser Gln Glu Leu Lys Asn Ser Ala Val Asn Leu 805 810 815
- Phe Asn Thr Thr Ala Ile Val Val Ala Glu Gly Thr Asp Arg Ile Ile 820 825 830

Glu Val Val Gln Arg Leu Cys Arg Ala Ile Leu His Ile Pro Arg Arg 835 840 845

Ile Arg Gln Gly Leu Glu Arg Phe Leu Leu * 850 855

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 854 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: not relevant
 - (ii) MOLECULE TYPE: protein
 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Arg Ala Arg Glu Lys Glu Arg Asn Cys Gln Asn Leu Trp Lys Trp 1 5 10 15

Gly Ile Met Leu Leu Gly Met Leu Met Thr Cys Ser Ala Ala Glu Asp 20 25 30

Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Ile Trp Lys Glu Ala Thr
35 40 45

Thr Thr Leu Phe Cys Ala Ser Asp Ala Lys Ala Tyr Lys Lys Glu Ala 50 55 60

His Asn Ile Trp Ala Thr His Ala Cys Val Pro Thr Asp Pro Asn Pro 65 70 75 80

Gln Glu Ile Glu Leu Glu Asn Val Thr Glu Asn Phe Asn Met Trp Lys 85 90 95

Asn Asn Met Val Glu Gln Met His Glu Asp Ile Ile Ser Leu Trp Asp 100 105 110

Gln Ser Leu Lys Pro Cys Val Lys Leu Thr Pro Leu Cys Val Thr Leu 115 120 125

Asn Cys Thr Asp Leu Lys Asn Glu Thr Lys Thr Asn Ser Ser Asp Ala 130 135 140

Asn Ser Asn Ser Gly Glu Ile Met Gly Asn Glu Glu Ile Lys Asn Cys 145 150 155 160

Ser Phe Asn Val Ser Thr Gly Ala Pro Gly Lys Val Gln Lys Glu Tyr 165 170 175

Ser Leu Phe Tyr Ala Leu Asp Asn Asn Asn Arg Thr Asn Ser Thr Asn 180 185 190

Tyr Arg Leu Ile Asn Cys Asp Thr Ser Thr Ile Thr Gln Ala Cys Pro

- Lys Ile Ser Phe Glu Pro Ile Pro Ile His Phe Cys Ala Pro Ala Gly 210 215 220
- Phe Ala Ile Leu Lys Cys Arg Asp Lys Lys Phe Asn Gly Thr Gly Pro 225 230 235 240
- Cys Ser Asn Val Ser Thr Val Gln Cys Thr His Gly Ile Arg Pro Val 245 250 255
- Val Ser Thr Gln Leu Leu Leu Asn Gly Ser Leu Ala Glu Glu Ile 260 265 270
- Ile Ile Arg Ser Glu Asn Leu Thr Asn Asn Val Lys Thr Ile Ile Val 275 280 285
- Gln Leu Asn Ala Ser Ile Val Ile Asn Cys Thr Arg Pro Tyr Lys Tyr 290 295 300
- Thr Arg Gln Arg Thr Ser Ile Gly Leu Arg Gln Ser Leu Tyr Thr Ile 305 310 315 320
- Thr Gly Lys Lys Lys Thr Gly Tyr Ile Gly Gln Ala His Cys Lys 325 330 335
- Ile Ser Arg Ala Glu Trp Asn Lys Ala Leu Gln Gln Val Ala Thr Lys
 340 345 350
- Leu Gly Asn Leu Leu Asn Lys Thr Thr Ile Thr Phe Lys Pro Ser Ser 355 360 365
- Gly Gly Asp Pro Glu Ile Thr Ser His Met Leu Asn Cys Gly Gly Asp 370 375 380
- Phe Phe Tyr Cys Asn Thr Ser Arg Leu Phe Asn Ser Thr Trp Asn Gln
- Thr Asn Ser Thr Gly Phe Asn Asn Gly Thr Val Thr Leu Pro Cys Arg
 405 410 415
- Ile Lys Gln Ile Val Asn Leu Trp Gln Arg Val Gly Lys Ala Met Tyr
 420 425 430
- Ala Pro Pro Ile Glu Gly Leu Ile Lys Cys Ser Ser Asn Ile Thr Gly
 435 440 445
- Leu Leu Thr Arg Asp Gly Gly Ala Asn Asn Ser Ser His Glu Thr 450 455 460
- Ile Arg Pro Gly Gly Gly Asp Met Arg Asp Asn Trp Arg Ser Glu Leu 465 470 475 480
- Tyr Lys Tyr Lys Val Val Lys Ile Glu Pro Ile Gly Val Ala Pro Thr 485 490 495

Lys Ala Arg Arg Arg Val Val Glu Arg Glu Lys Arg Ala Ile Gly Leu
500 505 510

- Gly Ala Val Phe Leu Gly Phe Leu Gly Ala Ala Gly Ser Thr Met Gly 515 520 525
- Ala Ala Ser Val Thr Leu Thr Val Gln Ala Arg Gln Leu Met Ser Gly 530 540
- Ile Val His Gln Gln Asn Asn Leu Leu Arg Ala Ile Glu Ala Gln Gln 545 550 555 560
- His Leu Gln Leu Thr Val Trp Gly Ile Lys Gln Leu Gln Ala Arg
 565 570 575
- Val Leu Ala Val Glu Arg Tyr Leu Arg Asp Gln Gln Leu Leu Gly Ile 580 585 590
- Trp Gly Cys Ser Gly Arg His Ile Cys Thr Thr Asn Val Pro Trp Asn 595 600 605
- Ser Ser Trp Ser Asn Arg Ser Leu Asp Glu Ile Trp Gln Asn Met Thr 610 615 620
- Trp Met Glu Trp Glu Arg Glu Ile Asp Asn Tyr Thr Gly Leu Ile Tyr 625 630 635 640
- Ser Leu Ile Glu Glu Ser Gln Ile Gln Gln Glu Lys Asn Glu Lys Glu 645 650 655
- Leu Leu Glu Leu Asp Lys Trp Ala Ser Leu Trp Asn Trp Phe Ser Ile 660 665 670
- Thr Lys Trp Leu Trp Tyr Ile Lys Leu Phe Ile Met Ile Val Gly Gly 675 680 685
- Leu Ile Gly Leu Arg Ile Val Phe Ala Val Leu Ser Val Val Asn Arg 690 695 700
- Val Arg Gln Gly Tyr Ser Pro Leu Ser Phe Gln Thr Leu Leu Pro Val 705 710 715 720
- Pro Arg Gly Pro Asp Arg Pro Glu Glu Ile Glu Glu Glu Gly Glu 725 730 735
- Arg Gly Arg Asp Arg Ser Ile Arg Leu Val Asn Gly Leu Phe Ala Leu 740 745 750
- Phe Trp Asp Asp Leu Arg Asn Leu Cys Leu Phe Ser Tyr His Arg Leu
 755 760 765
- Arg Asp Ser Ile Leu Ile Ala Ala Arg Ile Val Glu Leu Leu Gly Arg
 770 780
- Arg Gly Trp Glu Ala Leu Lys Tyr Leu Trp Asn Leu Leu Gln Tyr Trp 785 790 795 800

Ser Gln Glu Leu Arg Asn Ser Ala Ser Ser Leu Leu Asp Thr Ile Ala 805 810 815

Ile Ala Val Ala Glu Arg Thr Asp Arg Val Ile Glu Val Val Gln Arg 820 825 830

Ala Cys Arg Ala Ile Leu Asn Val Pro Arg Arg Ile Arg Gln Gly Leu 835 840 845

Glu Arg Leu Leu Leu Xaa 850

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 862 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: not relevant
- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Arg Val Lys Glu Lys Tyr Gln His Leu Trp Arg Trp Gly Trp Lys

1 10 15

Trp Gly Thr Met Leu Leu Gly Ile Leu Met Ile Cys Ser Ala Thr Glu 20 25 30

Lys Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Val Trp Lys Glu Ala 35 40 45

Thr Thr Leu Phe Cys Ala Ser Asp Ala Lys Ala Tyr Asp Thr Glu
50 55 60

Val His Asn Val Trp Ala Thr His Ala Cys Val Pro Thr Asp Pro Asn 65 70 75 80

Pro Gln Glu Val Val Leu Val Asn Val Thr Glu Asn Phe Asn Met Trp 85 90 95

Lys Asn Asp Met Val Glu Gln Met His Glu Asp Ile Ile Ser Leu Trp
100 105 110

Asp Gln Ser Leu Lys Pro Cys Val Lys Leu Thr Pro Leu Cys Val Ser 115 120 125

Leu Lys Cys Thr Asp Leu Gly Asn Ala Thr Asn Thr Asn Ser Ser Asn 130 135 140

Thr Asn Ser Ser Ser Gly Glu Met Met Met Glu Lys Gly Glu Ile Lys
145 150 155 160

Asn Cys Ser Phe Asn Ile Ser Thr Ser Ile Arg Gly Lys Val Gln Lys Glu Tyr Ala Phe Phe Tyr Lys Leu Asp Ile Ile Pro Ile Asp Asn Asp Thr Thr Ser Tyr Thr Leu Thr Ser Cys Asn Thr Ser Val Ile Thr Gln 200 Ala Cys Pro Lys Val Ser Phe Glu Pro Ile Pro Ile His Tyr Cys Ala Pro Ala Gly Phe Ala Ile Leu Lys Cys Asn Asn Lys Thr Phe Asn Gly 235 Thr Gly Pro Cys Thr Asn Val Ser Thr Val Gln Cys Thr His Gly Ile Arg Pro Val Val Ser Thr Gln Leu Leu Leu Asn Gly Ser Leu Ala Glu 265 Glu Glu Val Val Ile Arg Ser Ala Asn Phe Thr Asp Asn Ala Lys Thr Ile Ile Val Gln Leu Asn Gln Ser Val Glu Ile Asn Cys Thr Arg Pro Asn Asn Asn Thr Arg Lys Ser Ile Arg Ile Gln Arg Gly Pro Gly Arg 310 315 Ala Phe Val Thr Ile Gly Lys Ile Gly Asn Met Arg Gln Ala His Cys 330 Asn Ile Ser Arg Ala Lys Trp Asn Ala Thr Leu Lys Gln Ile Ala Ser Lys Leu Arg Glu Gln Phe Gly Asn Asn Lys Thr Ile Ile Phe Lys Gln 360 Ser Ser Gly Gly Asp Pro Glu Ile Val Thr His Ser Phe Asn Cys Gly 375 Gly Glu Phe Phe Tyr Cys Asn Ser Thr Gln Leu Phe Asn Ser Thr Trp 395 Phe Asn Ser Thr Trp Ser Thr Glu Gly Ser Asn Asn Thr Glu Gly Ser Asp Thr Ile Thr Leu Pro Cys Arg Ile Lys Gln Phe Ile Asn Met Trp

Gln Glu Val Gly Lys Ala Met Tyr Ala Pro Pro Ile Ser Gly Gln Ile

Arg Cys Ser Ser Asn Ile Thr Gly Leu Leu Thr Arg Asp Gly Gly

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Asn Asn Asn Gly Ser Glu Ile Phe Arg Pro Gly Gly Asp Met Arg Asp Asn Trp Arg Ser Glu Leu Tyr Lys Tyr Lys Val Val Lys Ile Glu Pro Leu Gly Val Ala Pro Thr Lys Ala Lys Arg Arg Val Val Gln 500 505 Arg Glu Lys Arg Ala Val Gly Ile Gly Ala Leu Phe Leu Gly Phe Leu Gly Ala Ala Gly Ser Thr Met Gly Ala Arg Ser Met Thr Leu Thr Val Gln Ala Arg Gln Leu Leu Ser Gly Ile Val Gln Gln Gln Asn Asn Leu Leu Arg Ala Ile Glu Ala Gln Gln His Leu Leu Gln Leu Thr Val Trp 565 570 Gly Ile Lys Gln Leu Gln Ala Arg Ile Leu Ala Val Glu Arg Tyr Leu 585 Lys Asp Gln Gln Leu Leu Gly Ile Trp Gly Cys Ser Gly Lys Leu Ile Cys Thr Thr Ala Val Pro Trp Asn Ala Ser Trp Ser Asn Lys Ser Leu Glu Gln Ile Trp Asn Asn Met Thr Trp Met Glu Trp Asp Arg Glu Ile 630 635 Asn Asn Tyr Thr Ser Leu Ile His Ser Leu Ile Glu Glu Ser Gln Asn Gln Gln Glu Lys Asn Glu Gln Glu Leu Leu Glu Leu Asp Lys Trp Ala 665 Ser Leu Trp Asn Trp Phe Asn Ile Thr Asn Trp Leu Trp Tyr Ile Lys Ile Phe Ile Met Ile Val Gly Gly Leu Val Gly Leu Arg Ile Val Phe 695 Ala Val Leu Ser Ile Val Asn Arg Val Arg Gln Gly Tyr Ser Pro Leu Ser Phe Gln Thr His Leu Pro Thr Pro Arg Gly Pro Asp Arg Pro Glu Gly Ile Glu Glu Glu Gly Gly Glu Arg Asp Arg Asp Arg Ser Ile Arg

Leu Val Asn Gly Ser Leu Ala Leu Ile Trp Asp Asp Leu Arg Ser Leu
755 760 765

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Cys Leu Phe Ser Tyr His Arg Leu Arg Asp Leu Leu Leu Ile Val Thr

Arg Ile Val Glu Leu Leu Gly Arg Arg Gly Trp Glu Ala Leu Lys Tyr 785 790 795 800

Trp Trp Asn Leu Leu Gln Tyr Trp Ser Gln Glu Leu Lys Asn Ser Ala 805 810 815

Val Ser Leu Leu Asn Ala Thr Ala Ile Ala Val Ala Glu Gly Thr Asp 820 825 830

Arg Val Ile Glu Val Val Gln Gly Ala Cys Arg Ala Ile Arg His Ile 835 840 845

Pro Arg Arg Ile Arg Gln Gly Leu Glu Arg Ile Leu Leu Xaa 850 855 860

(2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 856 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: not relevant
- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Lys Val Lys Gly Thr Arg Arg Asn Tyr Gln His Leu Trp Arg Trp 1 5 10 15

Gly Thr Leu Leu Gly Met Leu Met Ile Cys Ser Ala Thr Glu Lys 20 25 30

Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Val Trp Lys Glu Ala Thr 35 40 45

Thr Thr Leu Phe Cys Ala Ser Asp Ala Arg Ala Tyr Asp Thr Glu Val 50 55 60

His Asn Val Trp Ala Thr His Ala Cys Val Pro Thr Asp Pro Asn Pro 65 70 75 80

Gln Glu Val Val Leu Gly Asn Val Thr Glu Asn Phe Asn Met Trp Lys
85 90 95

Asn Asn Met Val Glu Gln Met Gln Glu Asp Ile Ile Ser Leu Trp Asp 100 105 110

Gln Ser Leu Lys Pro Cys Val Lys Leu Thr Pro Leu Cys Val Thr Leu 115 120 125

Asn Cys Thr Asp Leu Gly Lys Ala Thr Asn Thr Asn Ser Ser Asn Trp

- Lys Glu Glu Ile Lys Gly Glu Ile Lys Asn Cys Ser Phe Asn Ile Thr 145 150 155 160
- Thr Ser Ile Arg Asp Lys Ile Gln Lys Glu Asn Ala Leu Phe Arg Asn 165 170 175
- Leu Asp Val Val Pro Ile Asp Asn Ala Ser Thr Thr Thr Asn Tyr Thr
- Asn Tyr Arg Leu Ile His Cys Asn Arg Ser Val Ile Thr Gln Ala Cys 195 200 205
- Pro Lys Val Ser Phe Glu Pro Ile Pro Ile His Tyr Cys Thr Pro Ala 210 215 220
- Gly Phe Ala Ile Leu Lys Cys Asn Asn Lys Thr Phe Asn Gly Lys Gly
 225 230 235 240
- Pro Cys Thr Asn Val Ser Thr Val Gln Cys Thr His Gly Ile Arg Pro 245 250 255
- Ile Val Ser Thr Gln Leu Leu Leu Asn Gly Ser Leu Ala Glu Glu Glu 260 265 270
- Val Val Ile Arg Ser Asp Asn Phe Thr Asn Asn Ala Lys Thr Ile Ile 275 280 285
- Val Gln Leu Asn Glu Ser Val Ala Ile Asn Cys Thr Arg Pro Asn Asn 290 295 300
- Asn Thr Arg Lys Ser Ile Tyr Ile Gly Pro Gly Arg Ala Phe His Thr 305 310 315 320
- Thr Gly Arg Ile Ile Gly Asp Ile Arg Lys Ala His Cys Asn Ile Ser
- Arg Ala Gln Trp Asn Asn Thr Leu Glu Gln Ile Val Lys Lys Leu Arg
- Glu Gln Phe Gly Asn Asn Lys Thr Ile Val Phe Asn Gln Ser Ser Gly 355 360 365
- Gly Asp Pro Glu Ile Val Met His Ser Phe Asn Cys Arg Gly Glu Phe 370 380
- Phe Tyr Cys Asn Thr Thr Gln Leu Phe Asn Asn Thr Trp Arg Leu Asn 385 390 395 400
- His Thr Glu Gly Thr Lys Gly Asn Asp Thr Ile Ile Leu Pro Cys Arg
- Ile Lys Gln Ile Ile Asn Met Trp Gln Glu Val Gly Lys Ala Met Tyr
 420 425 430

Ala Pro Pro Ile Gly Gly Gln Ile Ser Cys Ser Ser Asn Ile Thr Gly
435 440 445

- Leu Leu Leu Thr Arg Asp Gly Gly Thr Asn Val Thr Asn Asp Thr Glu
 450 455 460
- Val Phe Arg Pro Gly Gly Gly Asp Met Arg Asp Asn Trp Arg Ser Glu 465 470 475 480
- Leu Tyr Lys Tyr Lys Val Ile Lys Ile Glu Pro Leu Gly Ile Ala Pro
 485 490 495
- Thr Lys Ala Lys Arg Arg Val Val Gln Arg Glu Lys Arg Ala Val Gly
 500 505 510
- Ile Val Gly Ala Met Phe Leu Gly Phe Leu Gly Ala Ala Gly Ser Thr 515 520 525
- Met Gly Ala Val Ser Leu Thr Leu Thr Val Gln Ala Arg Gln Leu Leu 530 535 540
- Ser Gly Ile Val Gln Gln Asn Asn Leu Leu Arg Ala Ile Glu Ala 545 550 555 560
- Gln Gln His Leu Leu Gln Leu Thr Val Trp Gly Ile Lys Gln Leu Gln 565 570 575
- Ala Arg Val Leu Ala Val Glu Arg Tyr Leu Arg Asp Gln Gln Leu Leu 580 585 590
- Gly Ile Trp Gly Cys Ser Gly Lys Leu Ile Cys Thr Thr Ala Val Pro 595 600 605
- Trp Asn Ala Ser Trp Ser Asn Lys Ser Leu Glu Asp Ile Trp Asp Asn 610 615 620
- Met Thr Trp Met Gln Trp Glu Arg Glu Ile Asp Asn Tyr Thr Asn Thr 625 630 . 635 640
- Ile Tyr Thr Leu Leu Glu Glu Ser Gln Asn Gln Glu Lys Asn Glu 645 650 655
- Gln Glu Leu Glu Leu Asp Lys Trp Ala Ser Leu Trp Asn Trp Phe 660 665 670
- Ser Ile Thr Asn Trp Leu Trp Tyr Ile Lys Ile Phe Ile Met Ile Val
- Gly Gly Leu Val Gly Leu Arg Ile Val Phe Ala Val Leu Ser Ile Val
 690 695 700
- Asn Arg Val Arg Gln Gly Tyr Ser Pro Leu Ser Phe Gln Thr Arg Leu 705 710 715 720
- Pro Val Pro Arg Gly Pro Asp Arg Pro Asp Gly Ile Glu Glu Gly 725 730 735

Gly Glu Arg Asp Arg Asp Arg Ser Val Arg Leu Val Asp Gly Phe Leu
740 745 750

- Ala Leu Ile Trp Glu Asp Leu Arg Ser Leu Cys Leu Phe Ser Tyr Arg
 755 760 765
- Arg Leu Arg Asp Leu Leu Leu Ile Ala Ala Arg Thr Val Glu Ile Leu
 770 780
- Gly His Arg Gly Trp Glu Ala Leu Lys Tyr Trp Trp Ser Leu Leu Gln 785 790 795 800
- Tyr Trp Ile Gln Glu Leu Lys Asn Ser Ala Val Ser Trp Leu Asn Ala 805 810 815
- Thr Ala Ile Ala Val Thr Glu Gly Thr Asp Arg Val Ile Glu Val Ala 820 825 830
- Gln Arg Ala Tyr Arg Ala Ile Leu His Ile His Arg Arg Ile Arg Gln 835 840 845
- Gly Leu Glu Arg Leu Leu Leu Xaa 850 855
- (2) INFORMATION FOR SEQ ID NO:6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 854 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: not relevant
 - (D) TOPOLOGY: not relevant
 - (ii) MOLECULE TYPE: protein
 - (iii) HYPOTHETICAL: NO
 - (ix) FEATURE:
 - (A) NAME/KEY: Region
 - (B) LOCATION: 1..854
 - (D) OTHER INFORMATION: /note= "Xaa indicates residues which are not specified in the consensus sequence."
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
 - Met Arg Xaa Xaa Glu Asn Xaa Xaa Xaa Xaa Xaa Xaa Xaa Trp Lys
 - Trp Gly Ile Met Leu Leu Gly Met Leu Met Kaa Ser Ala Kaa Glu Kaa 20 25 30
 - Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Xaa Lys Glu Thr Thr Thr
 35 40 45
 - Leu Phe Cys Ala Ser Asp Ala Lys Ala Tyr Xaa Xaa Glu Xaa His Asn 50 55 60

Xaa 65	Trp	Ala	Thr	His	Ala 70	Суз	Val	Pro	Thr	Asp 75	Pro	Asn	Pro	Gln	Glu 80
Ile	Xaa	Leu	Xaa	Asn 85	Val	Thr	Glu	Xaa	Phe 90	Asn	Met	Trp	Lys	Asn 95	Xaa
Met	Val	Glu	Gln 100	Met	His	Xaa	Asp	Ile 105	Ile	Ser	Leu	Trp	Asp 110	Xaa	Ser
Leu	Lys	Pro 115	Cys	Val	Lys	Leu	Thr 120	Pro	Leu	Сув	Val	Thr 125	Leu	Asn	Cys
Thr	Asp 130	Leu	Lys	Asn	Glu	Thr 135	Xaa	Xaa	Asn	Ser	Ser 140	Xaa	Xaa	Asn	Xaa
Xaa 145	Xaa	Xaa	Glu	Ile	Met 150	Gly	Xaa	Glu	Xaa	Xaa 155	Lys	Asn	Cys	Ser	Phe 160
Asn	Val	Xaa	Xaa	Xaa 165	Xaa	Xaa	Xaá	Xaa	Xaa 170	Xaa	Xaa	Xaa	Tyr	Xaa 175	Leu
Phe	Xaa	Xaa	Xaa 180	Xaa	Ile	Val	Xaa	Xaa 185	Xaa	Asn	Xaa	Xaa	Asn 190	Ser	Thr
Xaa	Xaa	Xaa 195	Leu	Xaa	Xaa	Cys	Xaa 200	Thr	Ser	Xaa	Xaa	Thr 205	Gln	Ala	Cys
Pro	Lys 210	Xaa	Ser	Phe	Glu	Pro 215	Ile	Pro	Ile	His	Xaa 220	Cys	Ala	Pro	Ala
Gly 225	Phe	Ala	Ile	Leu	Lys 230	Cys	Xaa	Asp	Lys	Lys 235	Phe	Asn	Gly	Thr	Gly 240
Pro	Cys	Xaa	Asn	Val 245	Ser	Thr	Val	Gln	Cys 250	Thr	His	Gly	Ile	Arg 255	Pro
Val	Val	Ser	Thr 260	Gln	Leu	Leu	Leu	Asn 265	Gly	Ser	Xaa	Ala	Glu 270	Glu	Glu
Xaa [.]	Xaa	Xaa 275	Arg	Ser	Xaa	Asn	Xaa 280	Xaa	Xaa	Asn	Xaa	Lys 285	Thr	Ile	Ile
Val	Gln 290	Leu	Asn	Xaa	Ser	Xaa 295	Xaa	Ile	Xaa	Cys	Thr 300	Arg	Pro	Xaa	Xaa
Xaa 305	Xaa	Xaa	Xaa	Xaa	Xaa 310	Xaa	Ile	Xaa	Xaa	Xaa 315	Xaa	Xaa	Xaa	Xaa	Xaa 320
Xaa	Thr	Gly	Lys	Xaa 325	Xaa	Lys	Xaa	Gly	Xaa 330	Xaa	Xaa	Gln	Ala	His 335	Cys
Xaa	Xaa	Xaa	Arg 340	Xaa	Xaa	Trp	Xaa	Xaa 345	Ala	Leu	Xaa	Gln	Xaa 350	Ala	Xaa
Lys	Leu	Xaa 355	Xaa	Xaa	Xaa		Lys 360	Thr	Xaa	Ile	Xaa	Phe 365	Xaa	Xaa	Ser

Ser Gly Gly Asp Xaa Glu Ile Xaa Xaa His Xaa Xaa Asn Cys Gly Gly Xaa Xaa Phe Tyr Cys Asn Thr Xaa Xaa Leu Phe Asn Ser Thr Trp Asn Xaa Thr Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Thr Leu Pro Cys 405 Arg Ile Xaa Gln Xaa Val Asn Xaa Trp Gln Xaa Val Gly Lys Ala Met 425 Tyr Ala Pro Pro Xaa Xaa Gly Xaa Ile Xaa Cys Ser Asn Ile Thr Gly Leu Leu Leu Thr Arg Asp Gly Gly His Asn Xaa Asn Asn Xaa Xaa Xaa Glu Thr Xaa Arg Pro Gly Xaa Gly Asp Met Arg Asp Asn Trp Arg Ser Glu Leu Tyr Lys Tyr Lys Val Xaa Lys Ile Glu Pro Xaa Gly Val Ala 490 Pro Thr Lys Ala Xaa Arg Arg Val Val Xaa Arg Glu Lys Arg Ala Xaa Met Gly Ala Xaa Phe Leu Gly Phe Leu Gly Ala Ala Gly Ser Thr Met Gly Ala Ala Ser Xaa Leu Thr Val Gln Ala Arg Gln Leu Xaa Ser Gly Ile Val Xaa Gln Gln Asn Asn Leu Leu Arg Ala Ile Glu Ala Gln Gln 550 555 His Leu Leu Gln Leu Thr Val Trp Gly Ile Lys Gln Leu Gln Ala Arg Val Leu Ala Val Glu Arg Tyr Leu Xaa Asp Gln Gln Leu Leu Gly Ile 580 585 Trp Gly Cys Ser Gly Xaa Xaa Ile Cys Thr Thr Xaa Val Pro Trp Asn 600 Xaa Ser Trp Ser Asn Xaa Ser Leu Asp Xaa Ile Trp Xaa Asn Met Thr 615 Trp Xaa Glu Trp Xaa Arg Glu Ile Xaa Asn Tyr Thr Xaa Leu Ile Xaa 635 Xaa Leu Ile Glu Glu Ser Gln Xaa Gln Gln Glu Lys Asn Glu Xaa Glu Leu Leu Glu Leu Asp Lys Trp Ala Ser Leu Trp Xaa Trp Phe Xaa Ile

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Xaa Xaa Trp Leu Trp Tyr Ile Lys Xaa Phe Ile Met Ile Val Xaa Gly 675 680 685

- Leu Xaa Gly Leu Arg Ile Val Phe Ala Val Leu Ser Xaa Val Asn Arg 690 695 700
- Val Arg Gln Gly Tyr Ser Pro Leu Ser Phe Gln Thr Xaa Xaa Pro Xaa 705 710 715 720
- Pro Arg Gly Pro Asp Arg Pro Xaa Xaa Ile Glu Xaa Glu Gly Glu 725 730 735
- Arg Xaa Arg Asp Arg Ser Xaa Arg Leu Val Xaa Gly Xaa Xaa Ala Leu 740 745 750
- Xaa Trp Xaa Asp Leu Arg Asn Leu Cys Leu Phe Ser Tyr His Arg Leu
 755 760 765
- Arg Asp Xaa Xaa Leu Ile Xaa Xaa Arg Ile Val Glu Leu Leu Gly Arg 770 775 780
- Arg Gly Trp Glu Ala Leu Lys Tyr Leu Trp Xaa Leu Leu Gln Tyr Trp 785 790 795 800
- Ser Gln Glu Leu Xaa Asn Ser Ala Xaa Xaa Leu Xaa Xaa Thr Xaa Ala 805 810 815
- Ile Xaa Val Ala Glu Xaa Thr Asp Arg Xaa Ile Glu Val Val Gln Arg 820 825 830
- Xaa Cys Arg Ala Ile Leu Xaa Xaa Pro Arg Arg Ile Arg Gln Gly Leu 835 840 845
- Glu Arg Xaa Leu Leu Xaa 850
- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 866 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: not relevant
 - (D) TOPOLOGY: not relevant
 - (ii) MOLECULE TYPE: protein
 - (iii) HYPOTHETICAL: NO
 - (ix) FEATURE:
 - (A) NAME/KEY: Region
 - (B) LOCATION: 1..866
 - (D) OTHER INFORMATION: /note= "Xaa residues are not specified in this consensus sequence."
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Met Arg Val Lys Glu Xaa Xaa Gln His Leu Trp Arg Trp Gly Trp Lys

1 5 10 15

- Trp Gly Xaa Met Leu Gly Xaa Leu Met Ile Cys Ser Ala Thr Glu 20 25 30
- Lys Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Val Trp Lys Glu Xaa 35 40 45
- Thr Thr Leu Phe Cys Ala Ser Asp Ala Lys Ala Tyr Xaa Xaa Glu 50 55 60
- Val His Asn Val Trp Ala Thr His Ala Cys Val Pro Thr Asp Pro Asn 65 70 75 80
- Pro Gln Glu Xaa Val Leu Xaa Asn Val Thr Glu Xaa Phe Asn Met Trp 85 90 95
- Lys Asn Xaa Met Val Glu Gln Met His Xaa Asp Ile Ile Ser Leu Trp
 100 105 110
- Asp Xaa Ser Leu Lys Pro Cys Val Lys Leu Thr Pro Leu Cys Val Xaa 115 120 125
- Leu Xaa Cys Thr Asp Leu Xaa Asn Xaa Thr Xaa Thr Asn Ser Ser Xaa 130 135 140
- Xaa Asn Ser Xaa Ser Gly Glu Xaa Met Met Xaa Xaa Xaa Glu Ile Lys 145 150 155 160
- Asn Cys Ser Phe Asn Xaa Ser Thr Xaa Xaa Xaa Gly Lys Val Gln Lys . 165 170 175
- Glu Tyr Xaa Xaa Phe Tyr Xaa Leu Asp Ile Val Ser Ile Xaa Xaa Xaa 180 185 190
- Asn Xaa Xaa Thr Ser Xaa Xaa Leu Thr Ser Cys Asn Thr Ser Val Xaa 195 200 205
- Thr Gln Ala Cys Pro Lys Val Ser Phe Glu Pro Ile Pro Ile His Tyr 210 215 220
- Cys Ala Pro Ala Gly Phe Ala Ile Leu Lys Cys Asn Xaa Lys Xaa Phe 225 230 235 240
- Asn Gly Thr Gly Pro Cys Xaa Asn Val Ser Thr Val Gln Cys Thr His 245 250 255
- Gly Ile Arg Pro Val Val Ser Thr Gln Leu Leu Leu Asn Gly Ser Xaa 260 265 270
- Ala Glu Glu Val Val Xaa Arg Ser Ala Asn Phe Xaa Asp Asn Ala 275 280 : 285
- Lys Thr Ile Ile Val Gln Leu Asn Xaa Ser Val Glu Ile Xaa Cys Thr 290 295 300

Arg Pro Asn Xaa Asn Xaa Xaa Lys Xaa Ile Arg Ile Xaa Arg Gly Xaa 305 310 315 320

- Gly Arg Xaa Phe Val Thr Xaa Xaa Lys Xaa Gly Xaa Xaa Xaa Gln Ala 325 330 335
- His Cys Xaa Xaa Xaa Arg Xaa Lys Trp Xaa Xaa Xaa Leu Lys Gln Ile 340 345 350
- Ala Ser Lys Leu Arg Glu Gln Phe Gly Asn Xaa Xaa Ile Ile Phe 355 360 365
- Xaa Xaa Ser Ser Gly Gly Asp Xaa Glu Ile Xaa Xaa His Ser Phe Asn 370 375 380
- Cys Gly Glu Xaa Phe Tyr Cys Asn Xaa Thr Xaa Leu Phe Asn Ser 385 390 395 400
- Thr Trp Phe Asn Xaa Thr Trp Ser Thr Glu Gly Ser Asn Asn Xaa Xaa 405 410. 415
- Gly Ser Xaa Xaa Ile Thr Leu Pro Cys Arg Ile Xaa Gln Phe Xaa Asn 420 425 430
- Met Trp Gln Xaa Val Gly Lys Ala Met Tyr Ala Pro Pro Xaa Xaa Gly
 435 440 445
- Gln Ile Arg Cys Xaa Ser Asn Ile Thr Gly Leu Leu Leu Thr Arg Asp 450 455 460
- Gly Gly His Asn Asp Asn Asn Xaa Xaa Xaa Glu Xaa Phe Arg Pro Gly 465 470 475 480
- Xaa Gly Asp Met Arg Asp Asn Trp Arg Ser Glu Leu Tyr Lys Tyr Lys
 485 490 495
- Val Xaa Lys Ile Glu Pro Leu Gly Val Ala Pro Thr Lys Ala Lys Arg
- Arg Val Val Gln Arg Glu Lys Arg Ala Val Gly Met Xaa Gly Ala Xaa
- Phe Leu Gly Phe Leu Gly Ala Ala Gly Ser Thr Met Gly Ala Xaa Ser 530 540
- Xaa Thr Leu Thr Val Gln Ala Arg Gln Leu Leu Ser Gly Ile Val Gln 545 550 560
- Gln Gln Asn Asn Leu Leu Arg Ala Ile Glu Ala Gln Gln His Leu Leu
 565 570 575
- Gln Leu Thr Val Trp Gly Ile Lys Gln Leu Gln Ala Arg Xaa Leu Ala 580 585 590
- Val Glu Arg Tyr Leu Lys Asp Gln Gln Leu Leu Gly Ile Trp Gly Cys 595 600 605

Ser Gly Lys Leu Ile Cys Thr Thr Xaa Val Pro Trp Asn Ala Ser Trp 610 615 620

Ser Asn Lys Ser Leu Xaa Gln Ile Trp Asn Asn Met Thr Trp Xaa Glu 625 630 635 640

Trp Asp Arg Glu Ile Xaa Asn Thr Xaa Leu Ile His Xaa Leu Ile Glu 645 650 655

Glu Ser Gln Asn Gln Gln Glu Lys Asn Glu Gln Glu Leu Leu Glu Leu
660 665 670

Asp Lys Trp Ala Ser Leu Trp Xaa Trp Phe Xaa Ile Xaa Asn Trp Leu 675 680 685

Trp Tyr Ile Lys Ile Phe Ile Met Ile Val Xaa Gly Leu Val Gly Leu 690 695 700

Arg Ile Val Phe Ala Val Leu Ser Ile Val Asn Arg Val Arg Gln Gly 705 710 715 720

Tyr Ser Pro Leu Ser Phe Gln Thr His Xaa Pro Xaa Pro Arg Gly Pro 725 730 735

Asp Arg Pro Xaa Gly Ile Glu Xaa Glu Gly Glu Arg Asp Arg Asp 740 745 750

Arg Ser Xaa Arg Leu Val Xaa Gly Xaa Leu Ala Leu Xaa Trp Xaa Asp 755 760 765

Leu Arg Xaa Leu Cys Leu Phe Ser Tyr His Arg Leu Arg Asp Leu Leu 770 780

Leu Ile Val Thr Arg Ile Val Glu Leu Leu Gly Arg Arg Gly Trp Glu 785 790 795 800

Ala Leu Lys Tyr Xaa Trp Xaa Leu Leu Gln Tyr Trp Ser Gln Glu Leu 805 810 815

Lys Asn Ser Ala Val Xaa Leu Xaa Asn Xaa Thr Ala Ile Xaa Val Ala 820 825 830

Glu Gly Thr Asp Arg Xaa Ile Glu Val Val Gln Xaa Xaa Cys Arg Ala 835 840 845

Ile Xaa His Ile Pro Arg Arg Ile Arg Gln Gly Leu Glu Arg Xaa Leu 850 860

Leu Xaa 865

(2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 836 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: not relevant

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- (D) TOPOLOGY: not relevant
- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: NO
- (ix) FEATURE:
 - (A) NAME/KEY: Region (B) LOCATION: 1..836
- (D) OTHER INFORMATION: /note= "Xaa residues are not specified in this consensus sequence."
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
 - Met Val Lys Glu Trp Trp Gly Leu Leu Gly Met Leu Met Ile Cys Ser 1 5 10 15
 - Ala Thr Glu Lys Leu Trp Val Thr Val Tyr Tyr Gly Val Pro Val Trp
 20 25 30
 - Lys Glu Thr Thr Leu Phe Cys Ala Ser Asp Ala Ala Tyr Glu Val
 - His Asn Val Trp Ala Thr His Ala Cys Val Pro Thr Asp Pro Asn Pro 50 55 60
 - Gln Glu Val Leu Asn Val Thr Glu Phe Asn Met Trp Lys Asn Met Val 65 70 75 80
 - Glu Gln Met Asp Ile Ile Ser Leu Trp Asp Ser Leu Lys Pro Cys Val 85 90 95
 - Lys Leu Thr Pro Leu Cys Val Thr Leu Asn Cys Thr Asp Leu Xaa Xaa 100 105 110
 - Xaa Thr Xaa Thr Asn Ser Ser Asp Ala Asn Xaa Xaa Xaa Xaa Xaa Xaa

 - Ile Xaa Xaa Xaa Xaa Xaa Xaa Xaa Asn Xaa Thr Xaa Xaa Xaa Leu Xaa
 165 170 175
 - Xaa Cys Asn Xaa Ser Val Xaa Thr Gln Ala Cys Pro Lys Val Ser Phe 180 185 190
 - Glu Pro Ile Pro Ile His Tyr Cys Xaa Pro Ala Gly Phe Ala Ile Leu 195 200 205
 - Lys Cys Asn Xaa Lys Xaa Phe Asn Gly Xaa Gly Pro Cys Xaa Asn Val 210 215 220

Ser Thr Val Gln Cys Thr His Gly Ile Arg Pro Xaa Val Ser Thr Gln 225 230 235 240

- Leu Leu Leu Asn Gly Ser Xaa Ala Glu Glu Glu Val Val Xaa Arg Ser 245 250 255
- Xaa Asn Phe Xaa Xaa Asn Ala Lys Thr Ile Ile Val Gln Leu Asn Xaa . 260 265 270
- Ser Val Xaa Ile Xaa Cys Thr Arg Pro Asn Xaa Asn Xaa Xaa Lys Xaa 275 280 285
- Ile Xaa Ile Xaa Xaa Gly Tyr Xaa Xaa Xaa Xaa Xaa Thr Xaa Arg Xaa 290 295 300
- Xaa Gly Asp Xaa Xaa Xaa Ala His Cys Xaa Xaa Xaa Arg Xaa Xaa Trp 305 310 315 320
- Xaa Asn Xaa Leu Xaa Gln Ile Xaa Xaa Lys Leu Arg Glu Gln Phe Gly
 325 330 335
- Asn Xaa Xaa Ile Xaa Phe Asn Xaa Ser Ser Gly Gly Asp Xaa Glu
 340 345 350
- Ile Xaa Met His Ser Phe Asn Cys Xaa Gly Glu Xaa Phe Tyr Cys Asn 355 360 365
- Thr Thr Xaa Leu Phe Asn Xaa Thr Trp Arg Xaa Xaa Thr Glu Xaa 370 375 380
- Xaa Xaa Xaa Xaa Xaa Ile Xaa Leu Pro Cys Arg Ile Xaa Gln Xaa 385 390 395 400
- Xaa Asn Met Trp Gln Xaa Val Gly Lys Ala Met Tyr Ala Pro Pro Xaa
 405 410 415
- Xaa Gly Gln Ile Xaa Cys Xaa Ser Asn Ile Thr Gly Leu Leu Thr 420 425 430
- Arg Asp Gly Gly Xaa Asn Xaa Asn Thr Xaa Xaa Glu Xaa Phe Arg 435 440 445
- Pro Gly Xaa Gly Asp Met Arg Asp Asn Trp Arg Ser Glu Leu Tyr Lys 450 455 460
- Tyr Lys Val Ile Lys Ile Glu Pro Leu Gly Xaa Ala Pro Thr Lys Ala 465 470 475 480
- Lys Arg Arg Val Val Gln Arg Glu Lys Arg Ala Val Gly Xaa Val Gly
 485 490 495
- Ala Met Phe Leu Gly Phe Leu Gly Ala Ala Gly Ser Thr Met Gly Ala 500 505 510
- Xaa Leu Thr Leu Thr Val Gln Ala Arg Gln Leu Leu Ser Gly Ile Val 515 520 525

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Gln Gln Gln Asn Asn Leu Leu Arg Ala Ile Glu Ala Gln Gln His Leu 530 535 540

- Leu Gln Leu Thr Val Trp Gly Ile Lys Gln Leu Gln Ala Arg Val Leu 545 550 555 560
- Ala Val Glu Arg Tyr Leu Xaa Asp Gln Gln Leu Leu Gly Ile Trp Gly 565 570 575
- Cys Ser Gly Lys Leu Ile Cys Thr Thr Xaa Val Pro Trp Asn Ala Ser 580 585 590
- Trp Ser Asn Lys Ser Leu Xaa Xaa Ile Trp Xaa Asn Met Thr Trp Xaa 595 600 605
- Xaa Trp Xaa Arg Glu Ile Xaa Asn Tyr Thr Asn Xaa Ile Xaa Xaa Leu 610 615 620
- Xaa Glu Glu Ser Gln Asn Gln Gln Glu Lys Asn Glu Gln Glu Leu Leu 625 630 635 640
- Glu Leu Asp Lys Trp Ala Ser Leu Trp Xaa Trp Phe Xaa Ile Xaa Asn
 645 650 . 655
- Trp Leu Trp Tyr Ile Lys Ile Phe Ile Met Ile Val Xaa Gly Leu Val
 660 665 670
- Gly Leu Arg Ile Val Phe Ala Val Leu Ser Ile Val Asn Arg Val Arg 675 680 685
- Gln Gly Tyr Ser Pro Leu Ser Phe Gln Thr Xaa Xaa Pro Xaa Pro Arg 690 695 700
- Gly Pro Asp Arg Pro Asp Gly Ile Glu Xaa Glu Gly Gly Glu Arg Asp 705 710 715 720
- Arg Asp Arg Ser Val Arg Leu Val Asp Gly Phe Leu Ala Leu Xaa Trp
 725 730 735
- Glu Asp Leu Arg Xaa Leu Cys Leu Phe Ser Tyr Xaa Arg Leu Arg Asp 740 745 750
- Leu Leu Leu Ile Xaa Xaa Arg Xaa Val Glu Xaa Leu Gly Xaa Arg Gly 755 760 765
- Trp Glu Ala Leu Lys Tyr Xaa Trp Ser Leu Leu Gln Tyr Trp Xaa Gln
 770 780
- Glu Leu Lys Asn Ser Ala Val Xaa Xaa Xaa Asn Xaa Thr Ala Ile Xaa 785 790 795 800
- Val Xaa Glu Gly Thr Asp Arg Xaa Ile Glu Val Xaa Gln Arg Xaa Xaa 805 810 815
- Arg Ala Ile Leu His Ile Xaa Arg Arg Ile Arg Gln Gly Leu Glu Arg 820 825 830

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Xaa Leu Leu Xaa 835

- (2) INFORMATION FOR SEQ ID NO:9:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 39 base pairs
 - (B) TYPE: nucleic acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "Oligonucleotide."
 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

CCCTTCGAAG AGGATATAAT CAGTTTATGG GATCAAAGC

39

- (2) INFORMATION FOR SEQ ID NO:10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 33 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "Oligonucleotide."
 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

CCCTTCGAAC TCTTCTTCTG CTAGACTGCC ATT

33

- (2) INFORMATION FOR SEQ ID NO:11:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 9193 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: not relevant
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (iv) ANTI-SENSE: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

GGTCTCTCTG GTTAGACCAG ATCTGAGCCT GGGAGCTCTC TGGCTAGCTA GGGAACCCAC

TGCTTAAGCC TCAATAAAGC TTGCCTTGAG TGCTTCAAGT AGTGTGTGCC CGTCTGTTGT 120

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GTGACTCTGG	TAACTAGAGA	TCCCTCAGAC	CCTTTTAGTC	AGTGTGGAAA	AATCTCTAGC	180
AGTGGCGCCC	GAACAGGGAC	CGGAAAGCAA	AAGGGAAACC	AGAGAAGCTC	TCTCGACGCA	240
GGACTCGGCT	TGCTGAAGCG	CGCACAGCAA	GAGGCGAGGG	GCGGCGACTG	GTGAGTACGC	300
CGATATTTT	GACTAGCGGA	GGCTAGAAGG	AGAGAGATGG	GTGCGAGAGC	GTCGGTATTA	360
AGCGGGGGAG	ATTTAGATCG	ATGGGAAAAA	ATTCGGTTAA	GGCCAGGGG	AAAGAAAAA	420
TATATGTTAA	AACATATAGT	ATGGGCAAGC	AGGGAGCTAG	AACGATTCGC	AGTCAATCCT	480
GGCCTGTTAG	AAACATCAGA	AGGCTGTAGA	CAAGTACTGG	AACAGCTACA	GCCATCCCTT	540
CAGACAGGAT	CAGAAGAACT	TAGATCCTTA	TATAATACAA	TAGCAACCCT	CTATGGTGTG	600
CATCAAAAAA	TAGAGGTAAA	AGACACCAAG	GAAGCTTTAG	ACAAAATAGA	GGAAGAGCAA	660
AACAAAAGTA	AGAAAAAAGC	ACAGCAAGCA	GCAGCTGACA	CAGGACACAG	CAGCCAGGTC	720
AGCCAAAATT	ACCCTATAGT	GCAGAACCTA	CAGGGGCAAA	TGGTACATCA	GGCCATATCA	780
CCTAGAACTT	TAAATGCATG	GGTAAAAGTA	ATAGAAGAGA	AGGCTTTCAG	CCCAGAAGTG	840
ATACCCATGT	TTTCAGCATT	ATCAGAAGGA	GCCACCCCAC	AAGATTTAAA	CACCATGTTA	900
AACACAGTGG	GGGGACATCA	AGCAGCCATG	CAAATGTTAA	AAGAGACCAT	CAATGAGGAA	960
GCTGCAGAAT	GGGATAGAGT	GCATCCAGTG	CAGGCAGGGC	TTATTGCACC	AGGCCAGATG	1020
AGAGAACCAA	GGGGAAGTGA	CATAGCAGGA	ACTACTAGTA	CCCTTCAGGA	ACAAATAGGA	1080
TGGATGACAA	GTAATCCACC	TATCCCAGTA	GGAGAAATTT	ATAAAAGGTG	GATAATCCTG	1140
GGCTTAAATA	AAATAGTAAG	AATGTATAGC	CCTATCAGCA	TTCTAGACAT	AAGACAAGGA	1200
CCAAAAGAAC	CCTTTAGAGA	CTATGTAGAC	CGGTTCTATA	AAACTCTAAG	AGCCGAGCAA	1260
GCTTCACAGG	AAGTAAAAA	TTGGATGACA	GAAACCTTGC	TGGTCCAAAA	TGCGAACCCA	1320
GATTGTAAGA	CTATCTTAAA	AGCATTAGGA	CCAGGAGCTA	CACTAGAAGA	AATGATGACA	1380
GCATGTCAGG	GAGTGGGAGG	ACCCGGCCAT	AAGGCAAGAG	TTTTAGCTGA	AGCAATGAGC	1440
CAAGTAACAA	ATTCAGCTGC	CATAATGATG	CAGAGAGGCA	ATTTTAAGAA	CCAAAGAAAG	1500
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AGAAAAAAGG	GCTGTTGGAA	ATGTGGAAAG	GAAGGACACC	AAATGAAAGA	TTGTATTGAG	1620
AGCCAGGCTA	ATTTTTTAGG	GAAAATCTGG	CCTTCCCACA	AGGGAAGGCC	AGGGAATTTC	1680
CTTCAGAGCA	GACCAGAGCC	AACAGCCCCA	CCAGAAGAGA	GCCTCAGGTC	TGGGATAGAG	1740
ACAACAACTC	CCTCTCAGAA	GCAGGAGCCA	ATAGACAAGG	AAGTGTATCC	TTTAACTTCC	1800
CTCAGATCAC	TCTTTGGCAA	CGACCCCTCG	TCACAATAAA	GATAGGGGGG	CAACTAAAGG	1860

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GATGGAAGCC	AAAAATGATA	GGGGGCATTG	GAGGTTTTAT	CAAAGTAAGA	CAGTATGATC	1980
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CTGTCAACAT	AATTGGAAGA	AATCTGTTGA	CTCAGATTGG	TTGCACTTTA	AATTTTCCCA	2100
TTAGTCCTAT	TGAAACTGTA	CCAGTGAGAT	TAAAGCCAGG	AATGGATGGC	CCAAAAGTTA	2160
AACAATGGCC	ATTGACAGAA	GAAAAAATAA	AAGCATTAGT	AGAAATTTGT	ACAGAAATGG	2220
AAAAGGAAGG	GAAAATTTCA	AAAATTGGGC	CTGAAAATCC	ATACAATACT	CCAGTATTTG	2280
CCATAAAGAA	AAAAGACAGT	ACTAAATGGA	GAAAATTAGT	AGATTTCAGA	GAACTTAATA	2340
AGAGAACTCA	AGACTTCTGG	GAAGTTCAAT	TAGGAATACC	ACATCCCTCA	GGTTTAAAAA	2400
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AAGATTTCAG	GAAGTATACT	ACATTTACCA	TACCTAGTAT	AAACAATCAG	ACACCAGGGA	2520
TTAGATATCA	GTACAATGTG	CTTCCACAGG	GATGGAAAGG	ATCACCAGCA	ATATTCCAAA	2580
GTAGCATGAC	AAAAATCTTA	GAGCCTTTTA	GAAAACAGAA	TCCAGACATA	GTTATCTATC	2640
AATACATGGA	TGATTTGTAT	GTAGGATCTG	ACTTAGAAAT	AGAGCAGCAT	AGAACAAAAA	2700
TAGAGGAACT	GAGACAGCAT	CTGTTGAGGT	GGGGATTTAC	CACACCAGAC	AAAAAACATC	2760
AGAAAGAACC	TCCATTCCTC	TGGATGGGGT	ATGAACTCCA	TCCTGATAAA	TGGACAGTAC	2820
AGCCTATAGT	GCTGCCAGAA	AAAGACAGCT	GGACTGTCAA	TGACATACAG	AAGTTAGTGG	2880
GAAAATTGAA	TTGGGCAAGT	CAGATTTACC	CAGGGATTAA	AGTAAGGCAA	TTATGTAAAC	2940
TCCTTAGAGG	AGCCAAAGCA	CTAACAGAAG	TAATACCACT	AACAGAAGGA	GCAGAGCTAG	3000
AACTGGCAGA	AAACAGAGAG	ATTCTAAAAG	AACCAGTACA	TGGAGTGTAT	TATGACCCAT	3060
CAAAAGACTT	AATAGCAGAA	ATACAGAAGC	AGGGGCAAGG	CCAATGGACA	TATCAAATTT	3120
ATCAAGAGCC	ATTTAAAAAT	CTGAAAACAG	GAAAATATGC	AAGAACGAGG	GGTGCCCACA	3180
CTAATGATGT	AAAACAATTA	ACAGAGGCAG	TGCAAAAAAT	AGCCACAGAA	AGCATAGTAA	3240
TATGGGGGAA	GACTCCTAAA	TTTAAACTGC	CCATACAAAA	GGAAACATGG	GAAACATGGT	3300
GGACAGAGTA	TTGGCAAGCC	ACTTGGATTC	CTGAATGGGA	GTTTGTCAAT	ACCCCTCCTT	3360
TAGTAAAATT	ATGGTACCAG	TTAGAGAAAG	AACCCATAGT	AGGAGCAGAA	ACTTTCTATG	3420
TAGATGGGGC	AGCTAACAGG	GAGACTAAAT	TAGGAAAAGC	AGGATATGTT	ACTAACAGAG	3480
GAAGACAAAA	GGTTGTCACC	CTGACTAACA	CAACAAATCA	GAAGACTGAG	TTACAAGCAA	3540
TTCATCTAGC	TTTACAGGAT	TCAGGATCAG	AAGTAAACAT	AGTAACAGAC	TCACAATATG	3600

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CAȚTAGGAAT	CATTCAAGCA	CAACCAGATC	AAAGTGAATC	AGAGTTAGTC	AATCAAATAA	3660			
TAGAGCAGTT	AATAAAAAAG	GAAAAGGTCT	ATCTGGCATG	GGTACCAGCA	CACAAAGGAA	3720			
TTGGAGGAAA	TGAACAAGTA	GATAAATTAG	TCAGTGCTGG	AATCAGAAAA	GTACTATTTT	3780			
TAGATGGAAT	AGATAAGGCC	CAAGAAGAAC	ATGAGAAATA	TCACAATAAT	TGGAGAGCAA	3840			

3900

5340 -

AATGTCAGCT AAAAGGAGAA GCTATGCATG GACAAGTAGA CTGTAGTCCA GGAATATGGC 3960

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AACTAGATTG TACACATTTA GAAGGAAAAG TTATCCTGGT AGCAGTTCAT GTAGCCAGTG 4020

GATATATAGA AGCAGAAGTT ATTCCAGCAG AAACAGGGCA GGAAACAGCA TACTTTCTCT 4080

TAAAATTAGC AGGAAGATGG CCAGTAAAAA CAATACATAC AGACAATGGC AGCAATTTCA 4140
CCAGTGCTGC GGTTAAGGCC GCCTGTTGGT GGGCAGGAGT CAAACAAGAA TTTGGAATTC 4200

CCTACAATCC CCAAAGTCAA GGAGTAGTAG AATCTATGAA TAAAGAATTA AAGAAAATTA 4260

TAGGACAGGT AAGAGATCAA GCTGAACATC TTAAGACAGC AGTACAAATG GCAGTATTTG 4320

TCCACAATTT TAAAAGAAAA GGGGGGATTG GGGGGTACAG TGCAGGGGAA AGAATAATAG 4380

ACATAATAGC AACAGACATA CAAACTAGAG AACTACAAAA ACAAATTACA AAAATTCAAA 4440

ATTTTCGGGT TTATTACAGG GACAGCAGAG ATCCACTTTG GAAAGGACCA GCAAAGCTCC 4500

TCTGGAAAGG TGAAGGGGCA GTAGTAATAC AAGATAATAG TGACATAAAA GTAGTGCCAA 4560

GAAGAAAAGC AAAGATCATT AGGGATTATG GAAAACAGAT GGCAGGTGAT GATTGTGTGG 4620

CAAGTAGACA GGATGAGGAT TAGAACATGG AAAAGTTTAG TAAAACATCA TGTATATGTT 4680

TCAAAGAAAG CTAGGGGATG GTTTTATAGA CATCACTATG AAAGCACTCA TCCAAAAATA 4740

AGTTCAGAAA TACACATCCC ACTAGGGGAT GCTAGATTGG TAGTAACAAC ATATTGGGGT 4800

CTGCATACAG GAGAAAGAGA ATGGCATTTG GGTCATGGAG TCTCCATAGA ATGGAGGAAA 4860

AGGAGCTATA GCACACAAGT AGACCCTGAA CTAGCAGACC AACTAATTCA TCTGTATTAC 4920
TTTGATTGTT TTTCAGACTC TGCTATAAGA AAGGCCTTAT TAGAACACAT AGTTAGCCCT 4980

AGGTGTGAAT ATCGAGCAGG ACATTCCAAG GTAGGATCTC TACAATACTT GGCACTATCA 5040

GCCTTAATAA CACCAAAGAA GATAAAGCCA CCTTTGCCTA GTGTTACGAA ACTGACAGAG 5100

GATAGATGGA ACAAGCCCCA GAAGACCAAG GGCCACAGAG GGAGCCATAC AATGAATGGA 5160

CACTAGAGCT TTTAGAGGAG CTTAAGGGAG AAGCTGTTAG ACATTTTCCT AGGCCATGGC 5220

TCCATAGCTT AGGACAACAT ATCTATGAAA CTTATGGAGA TACTTGGGCA GGAGTGGAAG 5280

CCATAATAAG AATTCTGCAA CAATTGCTGT TTATTCATTT CAGAATTGGG TGTCAACATA

- - WQ 98/5907	4				PCT/US	8/12990
GCAGAATAGG	CATTATTCAA	CAGAGGAGAG	CAAGAAGAAA	TGGAGCCAGT	AGATCCTAAT	5400
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AAAAGGTGTT	GCTTTCATTG	CCAAGTTTGT	TTCACAACAA	AAGCCTTAGG	CATCTCCTAT	5520
GGCAGGAAGA	AGCGGAGACA	GCGACGAAGA	CCTCGTCAGG	GCAGCCAGGC	TCATCAAGCT	5580
TCTCTATCAG	AGCAGTAAGT	AGTATATGTA	ATGCAACTTA	TATTAATTGT	AACAATAGTA	5640
GCTTTAGTAG	TAACATTAAT	AATAGCAATA	GTTGTGTGGT	CCATAGTACT	CATAGAATAT	5700
AGGAAAATAT	TAAGACAAAG	GAAAATAGAC	AAGTTAATTA	ATAGACTAGT	AGAAAGAGCA	5760
GAAGACAGTG	GCAATGAGAG	TGAAGGAGAA	CTGTCAGCAC	TTGTGGAGAT	GGGGGTGGAA	5820
ATGGGGCATC	ATGCTCCTTG	GGATGTTAAT	GATCTGTAGT	GCTACAGAAA	AATTGTGGGT	5880
CACAGTCTAT	TATGGGGTAC	CTGTGTGGAA	GGAAACAACT	ACCACTCTAT	TTTGTGCATC	5940
AGATGCTAAA	GCATATGAAG	AAGAGGTACA	TAATGTTTGG	GCCACACATG	CCTGTGTACC	6000
CACAGACCCC	AACCCACAAG	AAATAGTATT	GGCAAATGTG	ACAGAAGATT	TTAACATGTG	6060
GAAAAATGAA	ATGGTAGAAC	AGATGCATAC	TGATATAATC	AGTTTATGGG	ATGAAAGCCT	6120
AAAACCATGT	GTAAAATTAA	CCCCACTCTG	TGTTACTTTA	AATTGCACTG	ATTTGAAGAA	6180
TGAAACTAAG	ACCAATAGTA	GTGATGCCAA	TAGTAATAGC	GGGGAAATAA	TGGGGAACGA	6240
AGAGATAAAA	AATTGCTCTT	TCAATGTCAG	CACAGGCGCA	CCAGGTAAGG	TGCAGAAAGA	6300
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CCATATGTTG	ACAAGTTGTA	ACACCTCAGT	CAGTACACAG	GCCTGTCCAA	AGGTATCCTT	6420
TGAGCCAATT	CCCATACATT	ATTGTGCCCC	GGCTGGTTTT	GCAATTCTAA	AATGTAATGA	6480
TAAGAAGTTC	AATGGAACAG	GACCATGTAA	CAATGTCAGC	ACAGTACAAT	GTACACATGG	6540
AATTAGACCA	GTAGTGTCAA	CTCAACTGCT	GTTAAATGGC	AGTGTAGCAG	AAGAAGAGGT	6600
AGTACTTAGA	TCTGCCAATT	TCTCAGACAA	TGCTAAAACC	ATAATAGTAC	AGCTGAACCA	6660
CTCTGTAGAA	ATTACTTGTA	CAAGACCCAA	CTACAATGAA	ACAAAGAAAA	TCCGTATCCA	6720
CAGAGGATAT	GGAAGATCAT	TTGTTACAGT	AAGAAAATTG	GGAGATAGGA	AACAAGCACA	6780
TTGTACCATG	AATAGAACGA	AATGGGACAA	CGCTTTAAAA	CAGATAGCTA	GCAAATTAAG	6840
AGAACAATTT	ААТААТААА	CAGCAATAAT	CTTTAACCGG	TCCTCAGGAG	GGGACCTAGA	6900
AATTGAAATG	CACAGTTTTA	ATTGCGGAGG	GGAATTGTTC	TACTGTAATA	CAACAAAACT	6960
GTTTAATAGT	ACTTGGAATG	AGACTACAGA	GTCAAATGGC	AAGGGAGAAA	ATATCACACT	7020
CCCATGCAGA	ATAAGACAAT	TTGTAAACAT	GTGGCAGAAA	GTAGGAAAAG	CAATGTATGC	7080

CCCTCCCAGC	GATGGACAAA	TTAGGTGTAC	ATCAAATATT	ACTGGGCTAC	TATTAACAAG	7140
AGATGGGGGT	CATAATGATA	ACAACACTAA	CAACGAGACC	TTCAGACCGG	GAAGAGGAGA	7200
TATGAGGGAC	AATTGGAGAA	GTGAATTATA	тааататааа	GTAATAAAA	TTGAACCATT	7260
AGGAGTAGCA	CCCACCAAGG	CAAAGAGAAG	AGTGGTGCAG	AGAGAAAAA	GAGCAGTGGG	7320
AATGGTAGGA	GCTATGTTCC	TTGGGTTCTT	GGGAGCAGCA	GGAAGCACTA	TGGGCGCAGC	7380
GTCATTGACG	CTGACGGTAC	AGGCCAGACA	ATTATTGTCT	GGTATAGTGC	AGCAGCAGAA	7440
CAATCTGCTG	AGAGCTATTG	AGGCGCAACA	ACATCTGTTG	CAACTCACAG	TCTGGGGCAT	7500
CAAGCAGCTC	CAGGCAAGAG	TCCTGGCTGT	AGAAAGATAC	CTAAAGGATC	AACAGCTCCT	7560
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TTGGAGTAAT	AAATCTTTGG	ATCAGATTTG	GAATAACATG	ACCTGGATGG	AGTGGGACAG	7680
AGAAATTGCC	AATTACACAA	ACTTAATACA	TCACTTAATT	GAAGAATCGC	AAAACCAGCA	7740
AGAAAAGAAT	GAACAAGAAT	TATTGGAATT	AGATAAATGG	GCAAGTTTGT	GGAGTTGGTT	7800
TGACATATCA	AACTGGCTGT	GGTATATAAA	AATATTCATA	ATGATAGTAG	CAGGCTTAGT	7860
AGGTTTAAGA	ATAGTTTTTG	CTGTGCTTTC	TATAGTAAAT	AGAGTTAGGC	AGGGATACTC	7920
ACCATTGTCA	TTCCAGACCC	ACTTCCCAGC	TCCGAGGGGA	CCCGACAGGC	CAGACGGAAT	7980
CGAAGGAGAA	GGTGGAGAGA	GAGACAGAGA	CAGATCCGTG	CGATTAGTGG	ATGGATTCTT	8040
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CTTACTCTTG	ATTGTAACGA	GGATTGTGGA	ACTTCTCGGA	CGCAGGGGGT	GGGAAGCCCT	8160
CAAATATTTG	TGGAGTCTCC	TACAGTATTG	GAGTCAGGAG	CTAAAGAATA	GTGCTGTCAA	8220
CTTGTTCAAT	ACCACAGCTA	TAGTAGTAGC	TGAGGGGACA	GATAGGATCA	TAGAAGTAGT	8280
ACAAAGACTT	TGTAGAGCTA	TTCTCCACAT	ACCTAGAAGA	ATTAGACAGG	GCTTGGAAAG	8340
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TAAGGGAAAG	AATGAAACGA	GCTGGACCAG	CAGCAGATGG	GGTGGGAGCA	GCATCTCGAG	8460
ACCTAGAAAA	ACATGGAGCA	ATCACAAGTA	GCAATACAGC	AGCTACCAAT	GCTGATTGTG	8520
CCTGGCTAGA	AGCACAAGAG	GAGGAAGAGG	TGGGTTTTCC	AGTCAGACCT	CAGGTACCTT	8580
TAAGACCAAT	GACTTACAAG	GCAGGTATAG	ATCTTAGCCA	СТТТТТАААА	GAAAAGGGGG	8640
GACTGGAAGG	GCTAGTTTGG	TCCCAAAGAA	GACAAGATAT	CCTTGATCTG	TGGATCTACC	8700
ACACACAAGG	CTACTTCCCT	GATTGGCAGA	ACTACACACC	AGGGCCAGGG	ATCAGATATC	8760
CACTGACCTT	TGGATGGTGC	TTCAAGCTAG	TACCAGTTGA	GCCAGATAAG	GTAGAAGAGG	8820

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CCAATGAAGG	AGAGGACAAC	ATCTTGTTAC	ACCCTATGTG	TCTACATGGA	ATGGAGGACG	8880					
CGGAGAAAGA	AGTGTTAGTG	TGGAGATTTG	ACAGTAAACT	AGCCTTCCAT	CACGTAGCCC	8940					
GAGAGCTGCA	TCCGGAGTAC	TACAAAGACT	GCTGACATCG	AGCTTTCTAC	AAGGGACTTT	9000					
CCGCTGGGGA	CTTTCCAGGG	AGGCGTGATC	TGGGTGGGAC	TGGGGAGTGG	CGTGCCCTCA	9060					
GATGCTGCAT	ATAAGCAGCT	GCTTTTGCCT	GTACTGGGTC	TCTCTGGTTA	GACCAGATCT	9120					
GAGCCTGGGA	GCTCTCTGGC	TAGCTAGGGA	ACCCACTGCT	TAAGCCTCAA	TAAAGCTTGC	9180					
CTTGAGTGCT	TCA					9193					

(SEQ ID NO: 12)

HIVNCFULL7.SEQ tcaataaagcttgccttgagtgcttcaagtagtgtgtgcccgtctgttgtgtgactctggtaactagaga tccctcagacccttaaagtcagtgtggaaaaatctctagcagtggcgcccgaacagggacgcgaaagcga aagtagaaccagagaagctctctcgacgcaggactcggcttgctgaagcgcgcacagcaagaggcgaggg greggtartaagegggggagatttagategatgggaaaaaatteggttaaggecagggggaaagaaaaa tatatgttaaaacatataggtatgggcaagcagggagctagaacgattcgcagttaatcctggcctgttag aaacagctgaaggctgtagacaaatactagaacagctacagccatcccttcagacaggatcagaagaact tagatccttatatatatacaatagcaaccctctattgtgtgcatcaaaaaatagaggtaaaagacaccaag ctggcacaggaaacagcaggccaggtcagccaaaattaccctatagtgcagaacctacaggggcaaatggt acatcaggccatatcacctagaactttaaatgcatgggtaaaagtaatagaagagaaggctttcagccca gaagtgatacccatgttttcagcattatcagaaggagccaccccacaagatttaaacaccatgctaaaca cagrggggggacarcaagcagccargcaaargrtaaaagagaccarcaargaggaagcrgcagaatggga tagagtgcatccagtgcaggcagggcttattgcaccaggccagatgagagaaccaaggggaagtgacata aaatttataaaagatggataattttgggcttaaataaaatagtaagaatgtatagccctatcagcattct ggatataagacaaggaccaaaagaaccctttagagactatgtagaccggttctataaaactctaagagcc gagcaagcttcacaggaagtaaaaaattggatgacagaaaccttgttggtccaaaatgcgaacccagatt graagactattttaaaaagcattaggaccaggagctacattagaagaaatgatgacagcatgtcagggagt gggaggacccggccataaggcaagggttttagctgaagcaatgagccaagtaacaaattcagctgccata atgatgcagagaggcaattttaaaaaccaaagaaagatggttaagtgtttcaattgtggcaaagaagggc gaaagattgtattgaaagacaggctaattttttagggaaaatctggccttcccacaagggaaggccaggg aattttcttcagagcagaccagagccaacagccccaccagaagagagcctcaggtctgggatagagacaa caactccctctcagaagcaggagccaatagacaaggaagtgtatcctttaacttccctcagatcactctt tggcaacgacccctcgtcacaataaagataggggggcaactaaaggaagctctattagatacaggagcag atgatacagtattagaagaaatgaatttgccaggaagatggaaaccaaaaatgatagggggaattggagg tttatcaaagtaagacagtatgatcagatacccatagaaatctgtggacataaagctataggtacagta ttagtaggacctacacctgtcaacataattggaagaaatctgttgactcagattggttgcactttaaatt atttcaaaaattgggcctgaaaatccatacaatactccagtatttgccataaagaaaaagacagtacta aatggagaaaattagtagatttcagagaacttaataagagaactcaagacttctgggaagttcaattagg aataccacatcccgcagggttaaaaaagaaaaaatcagtaacagtactagatgtgggtgatgcatatttt tcagttcccttagatgaagatttcaggaagtatactgcatttaccatacctagtataaacaatgagacac cagggattagatatcagtacaatgtgctgccacagggatggaaaggatcaccagcaatattccaaagtag ttgtatgtaggatctgacttagaaatagagcagcatagaacaaaaatagaggaactgagacagcatctgt tgaagtggggatttaccacaccagacaaaaaacatcagaaggaacctccattcctctggatgggttatga actccatcctgataaatggacagtacagcctatagtgctgccagagaaagacagctggactgtcaatgac atacagaagttagtgggaaaattaaattgggcaagtcagatttacccagggattaaagtaaggcaattat gtaaactccttaggggagccaaagcactaacagatgtaataccactaacagaaggagcagagctagaact ggcagaaaacagagagattctaaaagaaccagtacatggagtgtattatgacccatcaaaagacttaata gcagaaatacagaagcagggacaaggtcaatggacatatcagatttatcaagagccatttaaaaatctga gaacaggaaaatatgcaagaacgaggggtgcccacactaatgatataaaacaattaacagaggcagtgca aaaaatagccacagaaagcatagtaatatggggaaagactcctaaatttaaactgcccatacaaaaggaa acatgggaaacatggtggacagagtattggcaagctacctggattcctgagtgggagtttgtcaataccc ctcctttagtaaaattatggtaccagttagagaaagaacccatagtaggagcagaaactttctatgtaga tggggcagctaacagggagactaaattaggaaaagcaggatatgttactaacagaggaagacaaaaggta gtccccctgactaacacaacaaatcagaagactgagttacaagcaattcatctagctttacaggattcag gatcagaagtaaacatagtaacagactcacaatatgcattaggaatcattcaagcacaaccagatcaaag tgaatcagagttagtcaatcaaataatagaacagttaataaaaaaggaaaaggtctatctggcatgggta ccagcacacaaaggaattggaggaaatgaacaagtagataaattagtcagtgctggaatcaggaaagtac tatttctagatggaatagataaggcccaagaagaacatgagaaatatcataattaggagagcaatggc tagtgattttaacctgccacctgtagtagcaaaggaaatagtagccagctgtgataaatgtcagctaaaa

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- WO 98/59074

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FOR SEQ ID NO:13 THROUGH SEQ ID NO:17, PLEASE SEE TABLE 5.

- (2) INFORMATION FOR SEQ ID NO:18
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "Oligonucleotide."
 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18

AAATCTCTAG CAGTGGCGCC CGAACAG

27

- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

GCACTCAAGG CAAGCTTTAT TGAGGCT

27

- (2) INFORMATION FOR SEQ ID NO:20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 31 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "Oligonucleotide."
 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

CACACACAG GCTACTTCCC TGATTGGCAG A

31

- (2) INFORMATION FOR SEQ ID NO:21:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 33 base pairs

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "Oligonucleotide."
- (iii) HYPOTHETICAL: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

ATGGAACAAG CCCCAGAAGA CCAAGGGCCA CAG

33

- (2) INFORMATION FOR SEQ ID NO:22:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 31 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

 - (iii) HYPOTHETICAL: NO
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

GGTCTGAGGG ATCTCTAGTT ACCAGAGTCA C

31

- (2) INFORMATION FOR SEQ ID NO:23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2404 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: not relevant
 - (ii) MOLECULE TYPE: protein
 - (iii) HYPOTHETICAL: YES
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

Met Glu Thr Ala Arg Gly Val Ala Leu Leu Tyr Ser Gly Leu Ala Ser 1 5 10 15

Asn Thr Tyr Arg Gly Leu Asn His Ile Ser Leu Glu Thr Arg Pro Ala 20 25 30

Arg Gly Thr Arg Pro Gly Leu Tyr Thr Arg Pro Leu Tyr Ser Thr Arg 35 40 45

Pro Gly Leu Tyr Ile Leu Glu Met Glu Thr Leu Glu Leu Glu Gly Leu 50 55 60

Tyr Met Glu Thr Leu Glu Met Glu Thr Ile Leu Glu Cys Tyr Ser Ser 65 70 75 80

- Glu Arg Ala Leu Ala Thr His Arg Gly Leu Leu Tyr Ser Leu Glu Thr 85 90 95
- Arg Pro Val Ala Leu Thr His Arg Val Ala Leu Thr Tyr Arg Thr Tyr
 100 105 110
- Arg Gly Leu Tyr Val Ala Leu Pro Arg Val Ala Leu Thr Arg Pro Leu 115 120 125
- Tyr Ser Gly Leu Ala Leu Ala Thr His Arg Thr His Arg Thr His Arg
- Leu Glu Pro His Glu Cys Tyr Ser Ala Leu Ala Ser Glu Arg Ala Ser 145 150 155 160
- Pro Ala Leu Ala Leu Tyr Ser Ala Leu Ala Thr Tyr Arg Gly Leu Gly
 165 170 . 175
- Leu Gly Leu Val Ala Leu His Ile Ser Ala Ser Asn Val Ala Leu Thr
 180 185 190
- Arg Pro Ala Leu Ala Thr His Arg His Ile Ser Ala Leu Ala Cys Tyr 195 200 205
- Ser Val Ala Leu Pro Arg Thr His Arg Ala Ser Pro Pro Arg Ala Ser 210 215 220
- Asn Pro Arg Gly Leu Asn Gly Leu Ile Leu Glu Val Ala Leu Leu Glu 225 230 235 240
- Ala Leu Ala Ala Ser Asn Val Ala Leu Thr His Arg Gly Leu Ala Ser
 245 250 255
- Pro Pro His Glu Ala Ser Asn Met Glu Thr Thr Arg Pro Leu Tyr Ser 260 265 270
- Ala Ser Asn Gly Leu Met Glu Thr Val Ala Leu Gly Leu Gly Leu Asn 275 280 285
- Met Glu Thr His Ile Ser Thr His Arg Ala Ser Pro Ile Leu Glu Ile 290 295 300
- Leu Glu Ser Glu Arg Leu Glu Thr Arg Pro Ala Ser Pro Gly Leu Ser 305 310 315 320
- Glu Arg Leu Glu Leu Tyr Ser Pro Arg Cys Tyr Ser Val Ala Leu Leu
 325 330 335
- Tyr Ser Leu Glu Thr His Arg Pro Arg Leu Glu Cys Tyr Ser Val Ala 340 345 350
- Leu Thr His Arg Leu Glu Ala Ser Asn Cys Tyr Ser Thr His Arg Ala 355 360 365

Ser	Pro 370	Leu	Glu	Leu	Tyr	Ser 375	Ala	Ser	Asn	Gly	Leu 380	Thr	His	Arg	Leu
Tyr 385	Ser	Thr	His	Arg	Ala 390	Ser	Asn	Ser	Glu	Arg 395	Ser	Glu	Arg	Ala	Ser 400
Pro	Ala	Leu	Ala	Ala 405	Ser	Asn	Ser	Glu	Arg 410	Ala	Ser	Asn	Ser	Glu 415	Arg
Gly	Leu	Tyr	Gly 420	Leu	Ile	Leu	Glu	Met 425	Glu	Thr	Gly	Leu	Tyr 430	Ala	Ser
Asn	Gly	Leu 435	Gly	Leu	Ile	Leu	Glu 440	Leu	Tyr	Ser	Ala	Ser 445	Asn	Cys	Tyr
Ser	Ser 450	Glu	Arg	Pro	His	Glu 455	Ala	Ser -	Asn	Val	Ala 460	Leu	Ser	Glu	Arg
Thr 465	His	Arg	Gly	Leu	Tyr 470	Ala	Leu	Ala		Arg 475	Gly	Leu	Tyr	Leu	Tyr 480
Ser	Val	Ala	Leu	Gly 485	Leu	Asn	Leu	Tyr	Ser 490	Gly	Leu	Thr	Tyr	Arg 495	Ser
Glu	Arg	Leu	Glu 500	Pro	His	Glu	Thr	Tyr 505	Arg	Ala	Leu	Ala	Leu 510	Glu	Ala
Ser	Pro	Ile 515	Leu	Glu	Val	Ala	Leu 520	Ser	Glu	Arg	Ile	Leu 525	Glu	Leu	Tyr
Ser	Ala 530	Ser	Asn	Gly	Leu	Ala 535	Ser	Asn	Ala	Ser	Asn 540	Ser	Glu	Arg	Thr
545	_		Glu		550					555					560
			Cys	565					570					575	
			Glu 580					585					590		
		595	Leu			•	600				_	605			
	610		Ile	٠.		615					620				
625			Ser		630			_		635		-			640
			Leu	645					650		-			655	
Ala	Ser	Asn	Ala 660	ser	Pro	Leu	Tyr	Ser 665	Leu	Tyr	Ser	Pro	His 670	GIU	Ala

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Ser Asn Gly Leu Tyr Thr His Arg Gly Leu Tyr Pro Arg Cys Tyr Ser 675 680 685

- Ala Ser Asn Ala Ser Asn Val Ala Leu Ser Glu Arg Thr His Arg Val 690 695 700
- Ala Leu Gly Leu Asn Cys Tyr Ser Thr His Arg His Ile Ser Gly Leu 705 710 715 720
- Tyr Ile Leu Glu Ala Arg Gly Pro Arg Val Ala Leu Val Ala Leu Ser 725 730 735
- Glu Arg Thr His Arg Gly Leu Asn Leu Glu Leu Glu Leu Glu Ala Ser
 740 745 750
- Asn Gly Leu Tyr Ser Glu Arg Leu Glu Ala Leu Ala Gly Leu Gly Leu
 755 760 765
- Gly Leu Val Ala Leu Val Ala Leu Leu Glu Ala Arg Gly Ser Glu Arg
 770 780
- Ala Leu Ala Ala Ser Asn Pro His Glu Ser Glu Arg Ala Ser Pro Ala 785 790 795 800
- Ser Asn Ala Leu Ala Leu Tyr Ser Thr His Arg Ile Leu Glu Ile Leu 805 810 815
- Glu Val Ala Leu Gly Leu Asn Leu Glu Ala Ser Asn His Ile Ser Ser 820 825 830
- Glu Arg Val Ala Leu Gly Leu Ile Leu Glu Thr His Arg Cys Tyr Ser 835 840 845
- Thr His Arg Ala Arg Gly Pro Arg Ala Ser Asn Thr Tyr Arg Ala Ser
- Asn Gly Leu Thr His Arg Leu Tyr Ser Leu Tyr Ser Ile Leu Glu Ala 865 870 875 880
- Arg Gly Ile Leu Glu His Ile Ser Ala Arg Gly Gly Leu Tyr Thr Tyr
 885 890 895
- Arg Gly Leu Tyr Ala Arg Gly Ser Glu Arg Pro His Glu Val Ala Leu 900 905 910
- Thr His Arg Val Ala Leu Ala Arg Gly Leu Tyr Ser Leu Glu Gly Leu 915 920 925
- Tyr Ala Ser Pro Ala Arg Gly Leu Tyr Ser Gly Leu Asn Ala Leu Ala 930 935 940
- His Ile Ser Cys Tyr Ser Thr His Arg Met Glu Thr Ala Ser Asn Ala 945 950 955 960
- Arg Gly Thr His Arg Leu Tyr Ser Thr Arg Pro Ala Ser Pro Ala Ser 965 970 975

Asn Ala Leu Ala Leu Glu Leu Tyr Ser Gly Leu Asn Ile Leu Glu Ala 980 985 990

- Leu Ala Ser Glu Arg Leu Tyr Ser Leu Glu Ala Arg Gly Gly Leu Gly 995 1000 1005
- Leu Asn Pro His Glu Ala Ser Asn Leu Tyr Ser Thr His Arg Ala Leu 1010 1015 1020
- Ala Ile Leu Glu Ile Leu Glu Pro His Glu Ala Ser Asn Ala Arg Gly 1025 1030 1035 1040
- Ser Glu Arg Ser Glu Arg Gly Leu Tyr Gly Leu Tyr Ala Ser Pro Leu 1045 1050 1055
- Glu Gly Leu Ile Leu Glu Gly Leu Met Glu Thr His Ile Ser Ser Glu 1060 1065 1070
- Arg Pro His Glu Ala Ser Asn Cys Tyr Ser Gly Leu Tyr Gly Leu Tyr 1075 1080 1085
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- Thr His Arg Thr His Arg Leu Tyr Ser Leu Glu Pro His Glu Ala Ser 1105 1110 1115 1120
- Asn Ser Glu Arg Thr His Arg Thr Arg Pro Ala Ser Asn Gly Leu Thr 1125 1130 1135
- His Arg Thr His Arg Gly Leu Ser Glu Arg Ala Ser Asn Gly Leu Tyr 1140 1145 1150
- Leu Tyr Ser Gly Leu Tyr Gly Leu Ala Ser Asn Ile Leu Glu Thr His
- Arg Leu Glu Pro Arg Cys Tyr Ser Ala Arg Gly Ile Leu Glu Ala Arg 1170 1175 1180
- Gly Gly Leu Asn Pro His Glu Val Ala Leu Ala Ser Asn Met Glu Thr 1185 1190 1195 1200
- Thr Arg Pro Gly Leu Asn Leu Tyr Ser Val Ala Leu Gly Leu Tyr Leu 1205 1210 1215
- Tyr Ser Ala Leu Ala Met Glu Thr Thr Tyr Arg Ala Leu Ala Pro Arg 1220 1225 1230
- Pro Arg Ser Glu Arg Ala Ser Pro Gly Leu Tyr Gly Leu Asn Ile Leu 1235 1240 1245
- Glu Ala Arg Gly Cys Tyr Ser Thr His Arg Ser Glu Arg Ala Ser Asn 1250 1255 1260
- Ile Leu Glu Thr His Arg Gly Leu Tyr Leu Glu Leu Glu Leu Glu Thr 1265 1270 1275 1280

His Arg Ala Arg Gly Ala Ser Pro Gly Leu Tyr Gly Leu Tyr Gly Leu
1285 1290 1295

- Tyr Pro Arg Ser Glu Arg Ala Ser Pro Ala Ser Asn Leu Tyr Ser Ala 1300 1305 1310
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- Leu Tyr Ser Val Ala Leu Ile Leu Glu Leu Tyr Ser Ile Leu Glu Gly
 1380 1385 1390
- Leu Pro Arg Leu Glu Gly Leu Tyr Val Ala Leu Ala Leu Ala Pro Arg 1395 1400 1405
- Thr His Arg Leu Tyr Ser Ala Leu Ala Leu Tyr Ser Ala Arg Gly Ala 1410 1415 1420
- Arg Gly Val Ala Leu Val Ala Leu Gly Leu Asn Ala Arg Gly Gly Leu 1425 1430 1435 1440
- Leu Tyr Ser Ala Arg Gly Ala Leu Ala Val Ala Leu Gly Leu Tyr Met 1445 1450 1455
- Glu Thr Val Ala Leu Gly Leu Tyr Ala Leu Ala Met Glu Thr Pro His 1460 1465 1470
- Glu Leu Glu Gly Leu Tyr Pro His Glu Leu Glu Gly Leu Tyr Ala Leu 1475 1480 1485
- Ala Ala Leu Ala Gly Leu Tyr Ser Glu Arg Thr His Arg Met Glu Thr 1490 1495 1500
- Gly Leu Tyr Ala Leu Ala Ala Leu Ala Ser Glu Arg Leu Glu Thr His 1505 1510 1515 1520
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- Arg Gly Gly Leu Asn Leu Glu Leu Glu Ser Glu Arg Gly Leu Tyr Ile 1540 1545 1550
- Leu Glu Val Ala Leu Gly Leu Asn Gly Leu Asn Gly Leu Asn Ala Ser 1555 1560 1565
- Asn Ala Ser Asn Leu Glu Leu Glu Ala Arg Gly Ala Leu Ala Ile Leu 1570 1575 1580

Glu Gly Leu Ala Leu Ala Gly Leu Asn Gly Leu Asn His Ile Ser Leu 1585 1590 1595 1600

- Glu Leu Glu Gly Leu Asn Leu Glu Thr His Arg Val Ala Leu Thr Arg 1605 1610 1615
- Pro Gly Leu Tyr Ile Leu Glu Leu Tyr Ser Gly Leu Asn Leu Glu Gly
 1620 1625 1630
- Leu Asn Ala Leu Ala Ala Arg Gly Val Ala Leu Leu Glu Ala Leu Ala 1635 1640 1645
- Val Ala Leu Gly Leu Ala Arg Gly Thr Tyr Arg Leu Glu Leu Tyr Ser 1650 1655 1660
- Ala Ser Pro Gly Leu Asn Gly Leu Asn Leu Glu Leu Glu Gly Leu Tyr 1665 1670 1675 1680
- Ile Leu Glu Thr Arg Pro Gly Leu Tyr Cys Tyr Ser Ser Glu Arg Gly
 1685 1690 1695
- Leu Tyr Leu Tyr Ser Leu Glu Ile Leu Glu Cys Tyr Ser Thr His Arg 1700 1705 1710
- Thr His Arg Thr His Arg Val Ala Leu Pro Arg Thr Arg Pro Ala Ser 1715 1720 1725
- Asn Ala Leu Ala Ser Glu Arg Thr Arg Pro Ser Glu Arg Ala Ser Asn 1730 1735 1740
- Leu Tyr Ser Ser Glu Arg Leu Glu Ala Ser Pro Gly Leu Asn Ile Leu 1745 1750 1755 1760
- Glu Thr Arg Pro Ala Ser Asn Ala Ser Asn Met Glu Thr Thr His Arg
- Thr Arg Pro Leu Glu Gly Leu Thr Arg Pro Ala Ser Pro Ala Arg Gly
 1780 1785 1790
- Gly Leu Ile Leu Glu Ala Leu Ala Ala Ser Asn Thr Tyr Arg Thr His 1795 1800 1805
- Arg Ala Ser Asn Leu Glu Ile Leu Glu His Ile Ser His Ile Ser Leu 1810 1815 1820
- Glu Ile Leu Glu Gly Leu Gly Leu Ser Glu Arg Gly Leu Asn Ala Ser 1825 1830 1835 1840
- Asn Gly Leu Asn Gly Leu Asn Gly Leu Leu Tyr Ser Ala Ser Asn Gly 1845 1850 1855
- Leu Gly Leu Asn Gly Leu Leu Glu Leu Glu Gly Leu Leu Glu Ala Ser 1860 1865 1870
- Pro Leu Tyr Ser Thr Arg Pro Ala Leu Ala Ser Glu Arg Leu Glu Thr 1875 1880 1885

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Arg Pro Ser Glu Arg Thr Arg Pro Pro His Glu Ala Ser Pro Ile Leu 1890 1895 1900

- Glu Ser Glu Arg Ala Ser Asn Thr Arg Pro Leu Glu Thr Arg Pro Thr 1905 1910 1915 1920
- Tyr Arg Ile Leu Glu Leu Tyr Ser Ile Leu Glu Pro His Glu Ile Leu 1925 1930 1935
- Glu Met Glu Thr Ile Leu Glu Val Ala Leu Ala Leu Ala Gly Leu Tyr 1940 1945 1950
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- Thr Tyr Arg His Ile Ser Ala Arg Gly Leu Glu Ala Arg Gly Ala Ser 2165 2170 2175
- Pro Leu Glu Leu Glu Leu Glu Ile Leu Glu Val Ala Leu Thr His Arg 2180 2185 2190

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Ala Arg Gly Ile Leu Glu Val Ala Leu Gly Leu Leu Glu Leu Glu Gly
2195 2200 2205

- Leu Tyr Ala Arg Gly Ala Arg Gly Gly Leu Tyr Thr Arg Pro Gly Leu 2210 2215 2220
- Ala Leu Ala Leu Glu Leu Tyr Ser Thr Tyr Arg Thr Arg Pro Thr Arg 2225 2230 2235 2240
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 2245 2250 2255
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- Asn Ser Glu Arg Ala Leu Ala Val Ala Leu Ala Ser Asn Leu Glu Pro 2275 2280 2285
- His Glu Ala Ser Asn Thr His Arg Ala Arg Gly Ala Leu Ala Ile Leu 2290 2295 2300
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- His Arg Ala Ser Pro Ala Arg Gly Ile Leu Glu Ile Leu Glu Gly Leu
 2325 2330 2335
- Val Ala Leu Val Ala Leu Gly Leu Asn Ala Arg Gly Leu Glu Cys Tyr 2340 2345 2350
- Ser Ala Arg Gly Ala Leu Ala Ile Leu Glu Leu Glu His Ile Ser Ile 2355 2360 2365
- Leu Glu Pro Arg Ala Arg Gly Ala Arg Gly Ile Leu Glu Ala Arg Gly 2370 2375 2380
- Gly Leu Asn Gly Leu Tyr Leu Glu Gly Leu Ala Arg Gly Pro His Glu 2385 2390 2395 2400

Leu Glu Leu Glu

We claim:

- An isolated human immunodeficiency virus type 1 (HIV-1), which is HIV-1_{JC}, identified by a nucleotide sequence as given in SEQ ID NO:11, or which is HIV-1_{NC}, identified by a nucleotide sequence as given in SEQ ID NO:12.
- 2. A biological sample comprising the HIV-1_{JC} of claim 1.
- 3. A biological sample comprising the HTV-1_{NC} of claim 1.
- 4. The biological sample of claim 1 wherein said sample is a sample of blood.
- 5. A biologically pure culture of host cells comprising the HIV- $1_{\rm JC}$ or HIV- $1_{\rm NC}$ of claim 1.
- 6. The culture of claim 5 wherein the host cells are peripheral blood mononuclear cells.
- 7. A composition comprising an antigenic preparation of the HIV- $1_{\rm JC}$ of claim 1.
- 8. A composition comprising an antigenic preparation of the HIV- 1_{NC} of claim 1.
- A kit for detecting the presence of HIV-1 antibodies comprising an antigenic preparation of the HIV-1_{JC} or HIV-1_{NC} of claim 1.
- An immunogenic composition comprising an antigenic preparation of the HIV-1_{JC} or HIV-1_{NC} of claim 1 and a pharmaceutically acceptable carrier.
- 11. A vaccine comprising an antigenic preparation of the HIV-1_{IC} or HIV-1_{NC} of claim 1.
- 12. An isolated DNA molecule comprising a nucleotide sequence encoding an infectious molecular clone for HIV-1_{JC} or an antigenic fragment thereof or for HIV-1_{NC} or an antigenic fragment thereof.

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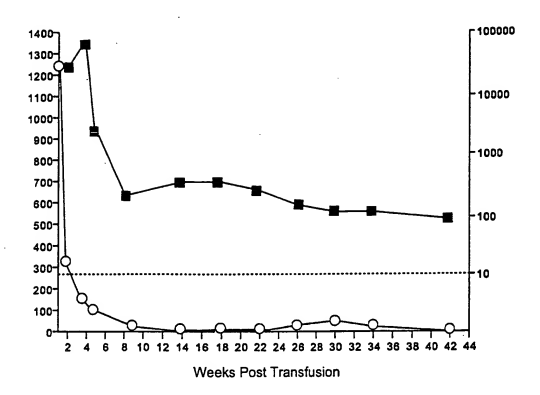
13. The isolated DNA molecule of claim 12, wherein the DNA comprises a nucleotide sequence encoding an HIV-1_{JC} envelope protein as given in SEQ ID NO:2, an HIV-1_{JC} gag, nef or p24 protein, or an antigenic fragment of any of the foregoing.

- 14. The isolated DNA molecule of claim 13, wherein the DNA comprises the nucleotide sequence of SEQ ID NO:1 or a portion thereof specifying an antigenic fragment of said HIV-1_{IC} envelope protein.
- 15. The isolated DNA molecule of claim 12, wherein the DNA comprises a nucleotide sequence encoding an HIV-1_{NC} envelope protein as given in SEQ ID NO:23, an HIV-1_{NC} gag, nef or p24 protein, or an antigenic fragment of any of the foregoing.
- 16. A method of inducing antibodies to HIV-1_{IC} in a nonhuman mammalian subject comprising the steps of (a) administering to the subject an immunogenic amount of an antigenic preparation of the HIV-1_{IC} and optionally (b) harvesting said antibodies to HIV-1_{IC}.
- 17. The method of claim 16 wherein said subject is a primate.
- 18. A method of inducing antibodies to HIV-1_{NC} in a nonhuman mammalian subject comprising the steps of (a) administering to the subject an immunogenic amount of an antigenic preparation of the HIV-1_{NC} and optionally (b) harvesting said antibodies to HIV-1_{NC}.
- 19. The method of claim 18 wherein said subject is a primate.
- 20. A method of immunizing a subject against the development of acquired immune deficiency syndrome (AIDS) comprising the step of administering to said subject an immunogenic amount of an antigenic preparation of HIV-1_{JC} or HIV-1_{NC} such that antibodies directed to HIV-1_{JC} or HIV-1_{NC} are produced in said subject and thereby symptoms of AIDS are diminished or prevented.
- 21. The method of claim 20 wherein said subject is a primate.

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22. A method for inducing acquired immune deficiency syndrome (AIDS) in a nonhuman primate, comprising the step of administering to said primate an effective amount of an infections preparation of the HIV-1 of claim 1 such that said primate develops AIDS.

- 23. The method of claim 22 wherein said infections preparation of HIV-1 is a biological specimen obtained from a nonhuman primate infected with HIV-1_{IC} and exhibiting AIDS.
- 24. The method of claim 22 wherein said infections preparation of HIV-1 is a biological specimen obtained from a nonhuman primate infected with HIV-1_{NC} and exhibiting AIDS.
- 25. A method of screening for HIV-1 in a biological sample comprising the step of introducing into said biological sample a hybridization probe comprising a nucleotide sequence comprising at least about 15 contiguous nucleotides from SEQ ID NO:11 or a sequence complementary thereto or from SEQ ID NO:12 or a nucleotide sequence complementary thereto under stringent conditions such that said hybridization probe binds to an HIV-1_{JC} gene or a nucleotide sequence having at least about 95% nucleotide sequence identity thereto or an HIV-1_{NC} gene or a nucleotide sequence having at least about 95% nucleotide sequence identity thereto.
- 26. A nonhuman primate model infected with at least one HIV-1 of claim 1 and exhibiting symptoms of AIDS, useful for the development of a drug or vaccine for the treatment or prevention of AIDS.
- 27. The primate model of claim 26 wherein said primate is a chimpanzee or a macaque.
- 28. The primate model of claim 27 wherein said HIV-1 is HIV-1_{jc}.
- 29. The primate model of claim 27 wherein said HIV-1 is HIV-1_{NC}.



■——■Plasma HIV-1 RNA (Equivalents, 10³/ml)

O——OAbsolute Circulating CD4* Cells/µl

FIG. 1

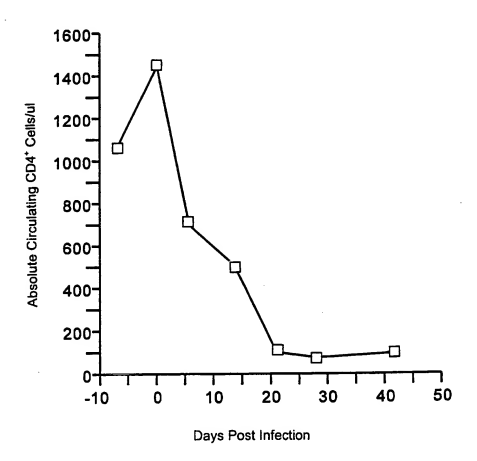
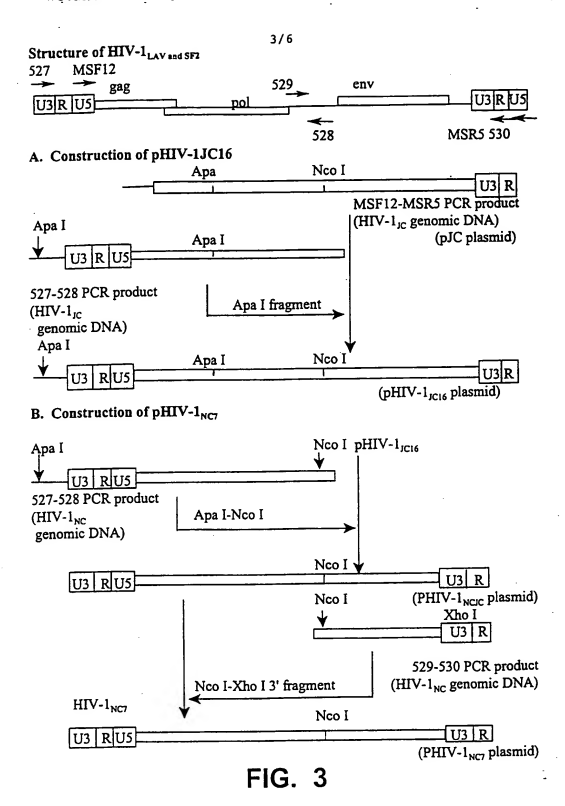


FIG. 2



SUBSTITUTE SHEET (RULE 26)



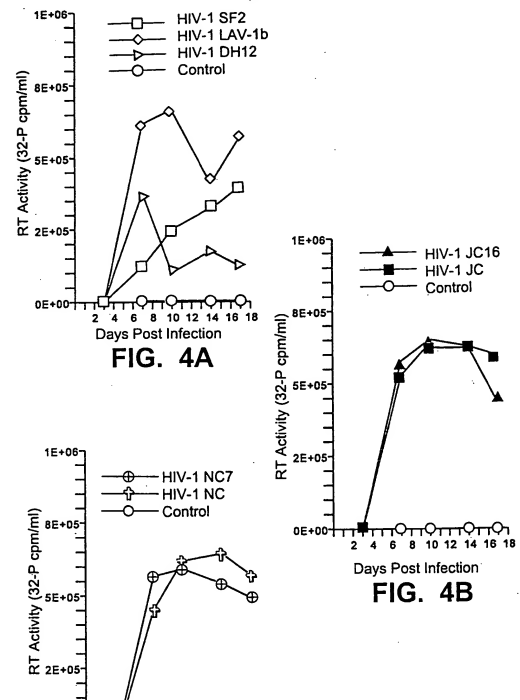


FIG. 4C

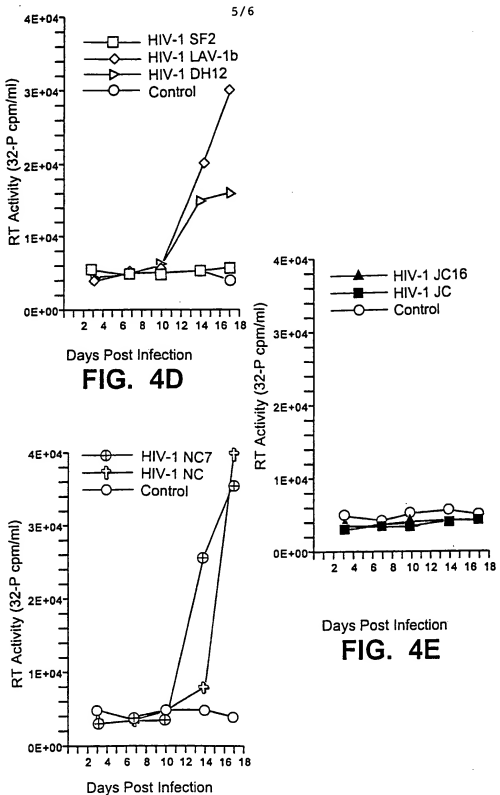
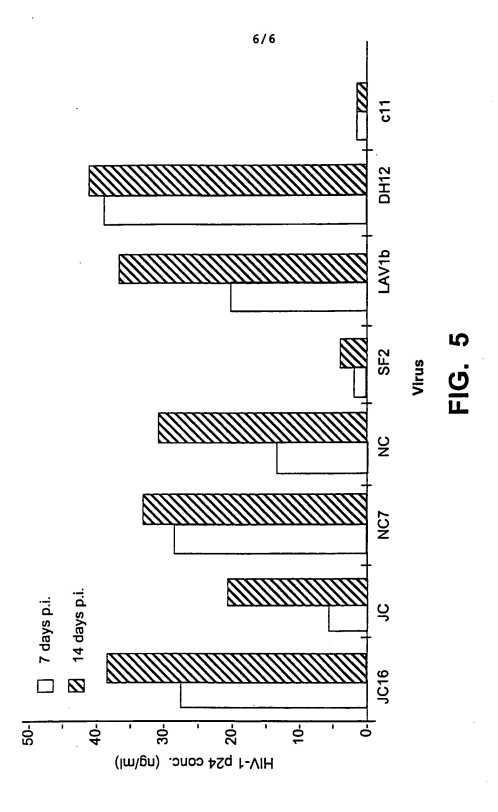


FIG. 4F

SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US98/12990

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :C12Q 1/68, 1/70; C12N 7/00, 7/01, 7/02; G01N 33/564; C07K 1/00 US CL : 435/5, 6, 235, 239, 974; 530/350, 826			
According to International Patent Classification (IPC) or to both national classification and IPC			
B. FIELDS SEARCHED			
Minimum documentation searched (classification system followed by classification symbols)			
U.S. : 435/5, 6, 235, 239, 974; 530/350, 826			
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched NONE			
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) APS, Medline, Aidsline			
C. DOCUMENTS CONSIDERED TO BE RELEVANT			
Category* Citation of document, with indication, where a	appropriate, of the relevant passages Relevant to claim No.		
NOVEMBRE et al Development of AIDS in a Chimpanzee Infected with Human Immunodeficiency Virus Type 1 Journal Of Virology. May 1997, Vol. 71, No. 5, pages 4086-4091, see entire document.			
VINCENT et al. Characterization of Resembling Human T-Cell Leukemia 1996, Vol. 226, No. 1, pages 57-65,	Virus Virology. 01 December		
Further documents are listed in the continuation of Box (C. See patent family annex.		
Special categories of cited documents:			
Special categories of cited documents: To later document published after the international filing data or priority data and not in conflict with the application but cited to understand to be of particular relevance.			
earlier document published on or efter the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step			
L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	when the document is taken alone 'Y' document of particular relevance; the claimed invention cannot be		
O* document referring to an oral disclosure, use, exhibition or other means	considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art		
document published prior to the international filing date but later than *&* document member of the same patent family			
Date of the actual completion of the international search 20 AUGUST 1998	Date of mailing of the international search report 18 SEP 1998		
lame and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	JEFFREY STUCKER TOWNER FOR		
acsimile No. (703) 305-3230	Tologham No. (702) 200 0106		

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/12990

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	
Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:	
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	╛
This International Searching Authority found multiple inventions in this international application, as follows:	1
Please See Extra Sheet.	
	l
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.	
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.	l
3. X As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 1-15 only to the extent of the elected specie, envelope protein.	
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims, it is covered by claims Nos.:	
Remark on Protest The additional search fees were accompanied by the applicant's protest.	
No protest accompanied the payment of additional search fees.	

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/12990

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claims 1-12, drawn to a virus.

Group II, claims 13-15, drawn to viral proteins.

Group III, claims 16-21, drawn to a method of inducing antibodies.

Group IV, claims 22-4 and 26-29, drawn to a primate model.

Group V, claim 25, drawn to a method of screening for HIV infection.

The inventions listed as Groups I-V do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: The rules of PCT unity of invention do not allow for multiple inventions. Each of the inventions requires a different search and a search of each of the inventions would not be co-extensive with each of the other searches.

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for more than one species to be searched, the appropriate additional search fees must be paid. The species are as follows:

Group II: envelope, gag, nef, and p24.

The claims are deemed to correspond to the species listed above in the following manner:

Group II, claims 13-15

The following claims are generic: 1-12.

The species listed above do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons: Each of the proteins is a different compound and has different characteristics such as biological function and immunological reactivity.